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In re Application of: ) Confirmation No. 7580  
 )  
Richard William Falla LE PAGE et al. ) Group Art Unit: 1645  
 )  
Application Number: 09/769,744 ) Examiner: DEVI  
 )  
Filed: January 26, 2001 )  
 )  
For: NUCLEIC ACIDS AND PROTEINS FROM STREPTOCOCCUS PNEUMONIAE

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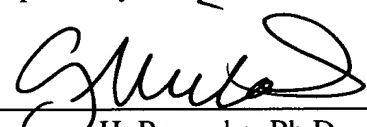
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Respectfully submitted,

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Microbial Techniques Limited  
Cortecs (UK) Limited 20 Trumpington Street  
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Lower Square  
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Middlesex TW7 6RL

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Patents ADP number (if you know it)

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If the applicant is a corporate body, give the country/state of its incorporation

4. Title of the invention

PROTEINS

5. Name of your agent (if you have one)

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## PROTEINS

The present invention relates to proteins derived from *Streptococcus pneumoniae*, nucleic acid molecules encoding such proteins, the use of the nucleic acid and/or proteins as antigens/immunogens and in detection/diagnosis, as well as methods for screening the proteins/nucleic acid sequences as potential anti-microbial targets.

5       *Streptococcus pneumoniae*, commonly referred to as the pneumococcus, is an important pathogenic organism. The continuing significance of *Streptococcus pneumoniae* infections in relation to human disease in developing and developed countries has been authoritatively reviewed (Fiber, G.R., *Science*, **265**: 1385-1387 (1994)). That indicates that on a global scale this organism is believed to be the most common bacterial cause of acute respiratory infections, and is estimated to result in 1 million childhood deaths each year, mostly in developing countries (Stansfield, S.K., *Pediatr. Infect. Dis.*, **6**: 622 (1987)). In the USA it has been suggested (Breiman *et al*, 10       *Arch. Intern. Med.*, **150**: 1401 (1990)) that the pneumococcus is still the most common cause of bacterial pneumonia, and that disease rates are particularly high in young children, in the elderly, and in patients with predisposing conditions such as asplenia, heart, lung and kidney disease, diabetes, alcoholism, or with immunosuppressive disorders, especially AIDS. These groups are at higher risk of pneumococcal 15       septicaemia and hence meningitis and therefore have a greater risk of dying from pneumococcal infection. The pneumococcus is also the leading cause of otitis media and sinusitis, which remain prevalent infections in children in developed countries, and which incur substantial costs.

20       The need for effective preventative strategies against pneumococcal infection is highlighted by the recent emergence of penicillin-resistant pneumococci. It has been reported that 6.6% of pneumococcal isolates in 13 US hospitals in 12 states were found to be resistant to penicillin and some isolates were also resistant to other antibiotics including third generation cyclosporins (Schappert, S.M., *Vital and Health Statistics of 25       the Centres for Disease Control/National Centre for Health Statistics*, **214**:1 (1992)).

The rates of penicillin resistance can be higher (up to 20%) in some hospitals (Breiman *et al*, J. Am. Med. Assoc., 271: 1831 (1994)). Since the development of penicillin resistance among pneumococci is both recent and sudden, coming after decades during which penicillin remained an effective treatment, these findings are  
5 regarded as alarming.

For the reasons given above, there are therefore compelling grounds for considering improvements in the means of preventing, controlling, diagnosing or treating pneumococcal diseases.

10 Various approaches have been taken in order to provide vaccines for the prevention of pneumococcal infections. Difficulties arise for instance in view of the variety of serotypes (at least 90) based on the structure of the polysaccharide capsule surrounding the organism. Vaccines against individual serotypes are not effective  
15 against other serotypes and this means that vaccines must include polysaccharide antigens from a whole range of serotypes in order to be effective in a majority of cases. An additional problem arises because it has been found that the capsular polysaccharides (each of which determines the serotype and is the major protective antigen) when purified and used as a vaccine do not reliably induce protective  
20 antibody responses in children under two years of age, the age group which suffers the highest incidence of invasive pneumococcal infection and meningitis.

A modification of the approach using capsule antigens relies on conjugating the polysaccharide to a protein in order to derive an enhanced immune response,  
25 particularly by giving the response T-cell dependent character. This approach has been used in the development of a vaccine against *Haemophilus influenzae*, for instance. There are, however, issues of cost concerning both the multi-polysaccharide vaccines and those based on conjugates.

30 A third approach is to look for other antigenic components which offer the potential to

be vaccine candidates. This is the basis of the present invention. Using a specially developed bacterial expression system, we have been able to identify a group of protein antigens from pneumococcus which are associated with the bacterial envelope or which are secreted.

5

Thus, in a first aspect the present invention provides a *Streptococcus pneumoniae* protein or polypeptide having a sequence selected from those shown in table 1.

10 In a second aspect, the present invention provides a *Streptococcus pneumoniae* protein or polypeptide having a sequence selected from those shown in table 2.

A protein or polypeptide of the present invention may be provided in substantially pure form. For example, it may be provided in a form which is substantially free of other proteins.

15

As discussed herein, the proteins and polypeptides of the invention are useful as antigenic material. Such material can be "antigenic" and/or "immunogenic". Generally, "antigenic" is taken to mean that the protein or polypeptide is capable of being used to raise antibodies or indeed is capable of inducing an antibody response in a subject.

20

"Immunogenic" is taken to mean that the protein or polypeptide is capable of eliciting a protective immune response in a subject. Thus, in the latter case, the protein or polypeptide may be capable of not only generating an antibody response but, in addition, a non-antibody based immune response.

25

The skilled person will appreciate that homologues or derivatives of the proteins or polypeptides of the invention will also find use in the context of the present invention, ie as antigenic/immunogenic material. Thus, for instance proteins or polypeptides which include one or more additions, deletions, substitutions or the like are encompassed by the present invention. In addition, it may be possible to replace one amino acid with another of similar "type". For instance replacing one hydrophobic amino acid with another.

30

One can use a program such as the CLUSTAL program to compare amino acid sequences. This program compares amino acid sequences and finds the optimal alignment by inserting spaces in either sequence as appropriate. It is possible to calculate amino acid identity or similarity (identity plus conservation of amino acid type) for an optimal alignment. A program like BLASTx will align the longest stretch of similar sequences and assign a value to the fit. It is thus possible to obtain a comparison where several regions of similarity are found, each having a different score. Both types of identity analysis are contemplated in the present invention.

In the case of homologues and derivatives, the degree of identity with a protein or polypeptide as described herein is less important than that the homologue or derivative should retain the antigenicity or immunogenicity of the original protein or polypeptide. However, suitably, homologues or derivatives having at least 60% similarity (as discussed above) with the proteins or polypeptides described herein are provided. Preferably, homologues or derivatives having at least 70% similarity, more preferably at least 80% similarity are provided. Most preferably, homologues or derivatives having at least 90% or even 95% similarity are provided.

In an alternative approach, the homologues or derivatives could be fusion proteins, incorporating moieties which render purification easier, for example by effectively tagging the desired protein or polypeptide. It may be necessary to remove the "tag" or it may be the case that the fusion protein itself retains sufficient antigenicity to be useful.

In an additional aspect of the invention there are provided antigenic/immunogenic fragments of the proteins or polypeptides of the invention, or of homologues or derivatives thereof.

For fragments of the proteins or polypeptides described herein, or of homologues or derivatives thereof, the situation is slightly different. It is well known that is possible to screen an antigenic protein or polypeptide to identify epitopic regions, ie those regions

which are responsible for the protein or polypeptide's antigenicity or immunogenicity. Methods for carrying out such screening are well known in the art. Thus, the fragments of the present invention should include one or more such epitopic regions or be sufficiently similar to such regions to retain their antigenic/immunogenic properties.

5 Thus, for fragments according to the present invention the degree of identity is perhaps irrelevant, since they may be 100% identical to a particular part of a protein or polypeptide, homologue or derivative as described herein. The key issue, once again, is that the fragment retains the antigenic/immunogenic properties.

10 Thus, what is important for homologues, derivatives and fragments is that they possess at least a degree of the antigenicity/immunogenicity of the protein or polypeptide from which they are derived.

15 Gene cloning techniques may be used to provide a protein of the invention in substantially pure form. These techniques are disclosed, for example, in J. Sambrook *et al Molecular Cloning* 2nd Edition, Cold Spring Harbor Laboratory Press (1989). Thus, in a third aspect, the present invention provides a nucleic acid molecule comprising or consisting of a sequence which is:

20 (i) any of the DNA sequences set out in Table 1 or their RNA equivalents;

(ii) a sequence which is complementary to any of the sequences of (i);

25 (iii) a sequence which codes for the same protein or polypeptide, as those sequences of (i) or (ii);

(iv) a sequence which has substantial identity with any of those of (i), (ii) and (iii);

- (v) a sequence which codes for a homologue, derivative or fragment of a protein as defined in Table 1.

5 In a fourth aspect the present invention provides a nucleic acid molecule comprising or consisting of a sequence which is:

- (i) any of the DNA sequences set out in Table 2 or their RNA equivalents;
- (ii) a sequence which is complementary to any of the sequences of (i);
- 10 (iii) a sequence which codes for the same protein or polypeptide, as those sequences of (i) or (ii);
- (iv) a sequence which has substantial identity with any of those of (i), (ii) and  
15 (iii); or
- (v) a sequence which codes for a homologue, derivative or fragment of a protein as defined in Table 2.

20 The nucleic acid molecules of the invention may include a plurality of such sequences, and/or fragments. The skilled person will appreciate that the present invention can include novel variants of those particular novel nucleic acid molecules which are exemplified herein. Such variants are encompassed by the present invention. These may occur in nature, for example because of strain variation. For example, additions,  
25 substitutions and/or deletions are included. In addition, and particularly when utilising microbial expression systems, one may wish to engineer the nucleic acid sequence by making use of known preferred codon usage in the particular organism being used for expression. Thus, synthetic or non-naturally occurring variants are also included within the scope of the invention.

The term "RNA equivalent" when used above indicates that a given RNA molecule has a sequence which is complementary to that of a given DNA molecule (allowing for the fact that in RNA "U" replaces "T" in the genetic code).

5 When comparing nucleic acid sequences for the purposes of determining the degree of homology or identity one can use programs such as BESTFIT and GAP (both from the Wisconsin Genetics Computer Group (GCG) software package) BESTFIT, for example, compares two sequences and produces an optimal alignment of the most similar segments. GAP enables sequences to be aligned along their whole length and finds the  
10 optimal alignment by inserting spaces in either sequence as appropriate. Suitably, in the context of the present invention when discussing identity of nucleic acid sequences, the comparison is made by alignment of the sequences along their whole length.

15 Preferably, sequences which have substantial identity have at least 50% sequence identity, desirably at least 75% sequence identity and more desirably at least 90 or at least 95% sequence identity with said sequences. In some cases the sequence identity may be 99% or above.

20 Desirably, the term "substantial identity" indicates that said sequence has a greater degree of identity with any of the sequences described herein than with prior art nucleic acid sequences.

25 It should however be noted that where a nucleic acid sequence of the present invention codes for at least part of a novel gene product the present invention includes within its scope all possible sequence coding for the gene product or for a novel part thereof.

The nucleic acid molecule may be in isolated or recombinant form. It may be incorporated into a vector and the vector may be incorporated into a host. Such vectors and suitable hosts form yet further aspects of the present invention.

Therefore, for example, by using probes based upon the nucleic acid sequences provided herein, genes in *Streptococcus pneumoniae* can be identified. They can then be excised using restriction enzymes and cloned into a vector. The vector can be introduced into a suitable host for expression.

5

Nucleic acid molecules of the present invention may be obtained from *S.pneumoniae* by the use of appropriate probes complementary to part of the sequences of the nucleic acid molecules. Restriction enzymes or sonication techniques can be used to obtain appropriately sized fragments for probing.

10

Alternatively PCR techniques may be used to amplify a desired nucleic acid sequence. Thus the sequence data provided herein can be used to design two primers for use in PCR so that a desired sequence, including whole genes or fragments thereof, can be targeted and then amplified to a high degree.

15

Typically primers will be at least 15-25 nucleotides long.

20

As a further alternative chemical synthesis may be used. This may be automated. Relatively short sequences may be chemically synthesised and ligated together to provide a longer sequence.

25

There is another group of proteins from *S.pneumoniae* which have been identified using the bacterial expression system described herein. These are known proteins from *S.pneumoniae*, which have not previously been identified as antigenic proteins. The amino acid sequences of this group of proteins, together with DNA sequences coding for them are shown in Table 3. These proteins, or homologues, derivatives and/or fragments thereof also find use as antigens/immunogens. Thus, in another aspect the present invention provides the use of a protein or polypeptide having a sequence selected from those shown in Tables 1-3, or homologues, derivatives and/or fragments thereof, as an immunogen/antigen.

30

In yet a further aspect the present invention provides an immunogenic/antigenic composition comprising one or more proteins or polypeptides selected from those whose sequences are shown in Tables 1-3, or homologues or derivatives thereof, and/or fragments of any of these. In preferred embodiments, the immunogenic/antigenic composition is a vaccine or is for use in a diagnostic assay.

In the case of vaccines suitable additional excipients, diluents, adjuvants or the like may be included. Numerous examples of these are well known in the art.

It is also possible to utilise the nucleic acid sequences shown in Tables 1-3 in the preparation of so-called DNA vaccines. Thus, the invention also provides a vaccine composition comprising one or more nucleic acid sequences as defined herein. DNA vaccines are described in the art (see for instance, Donnelly *et al*, *Ann. Rev. Immunol.*, 15:617-648 (1997)) and the skilled person can use such art described techniques to produce and use DNA vaccines according to the present invention.

As already discussed herein the proteins or polypeptides described herein, their homologues or derivatives, and/or fragments of any of these, can be used in methods of detecting/diagnosing *S.pneumoniae*. Such methods can be based on the detection of antibodies against such proteins which may be present in a subject. Therefore the present invention provides a method for the detection/diagnosis of *S.pneumoniae* which comprises the step of bringing into contact a sample to be tested with at least one protein, or homologue, derivative or fragment thereof, as described herein. Suitably, the sample is a biological sample, such as a tissue sample or a sample of blood or saliva obtained from a subject to be tested.

In an alternative approach, the proteins described herein, or homologues, derivatives and/or fragments thereof, can be used to raise antibodies, which in turn can be used to detect the antigens, and hence *S.pneumoniae*. Such antibodies form another aspect of

the invention. Antibodies within the scope of the present invention may be monoclonal or polyclonal.

5 Polyclonal antibodies can be raised by stimulating their production in a suitable animal host (e.g. a mouse, rat, guinea pig, rabbit, sheep, goat or monkey) when a protein as described herein, or a homologue, derivative or fragment thereof, is injected into the animal. If desired, an adjuvant may be administered together with the protein. Well-known adjuvants include Freund's adjuvant (complete and incomplete) and aluminium hydroxide. The antibodies can then be purified by virtue of their binding to a protein as  
10 described herein.

Monoclonal antibodies can be produced from hybridomas. These can be formed by fusing myeloma cells and spleen cells which produce the desired antibody in order to form an immortal cell line. Thus the well-known Kohler & Milstein technique (*Nature* 15 256 (1975)) or subsequent variations upon this technique can be used.

Techniques for producing monoclonal and polyclonal antibodies that bind to a particular polypeptide/protein are now well developed in the art. They are discussed in standard immunology textbooks, for example in Roitt *et al*, *Immunology* second edition (1989),  
20 Churchill Livingstone, London.

In addition to whole antibodies, the present invention includes derivatives thereof which are capable of binding to proteins etc as described herein. Thus the present invention includes antibody fragments and synthetic constructs. Examples of antibody fragments and synthetic constructs are given by Dougall *et al* in *Tibtech* 12 372-379 (September 25 1994).

Antibody fragments include, for example, Fab, F(ab')<sub>2</sub> and Fv fragments. Fab fragments (These are discussed in Roitt *et al* [*supra*]). Fv fragments can be modified to produce a synthetic construct known as a single chain Fv (scFv) molecule. This includes a peptide  
30

linker covalently joining  $V_h$  and  $V_l$  regions, which contributes to the stability of the molecule. Other synthetic constructs that can be used include CDR peptides. These are synthetic peptides comprising antigen-binding determinants. Peptide mimetics may also be used. These molecules are usually conformationally restricted organic rings that mimic the structure of a CDR loop and that include antigen-interactive side chains.

Synthetic constructs include chimaeric molecules. Thus, for example, humanised (or primatised) antibodies or derivatives thereof are within the scope of the present invention. An example of a humanised antibody is an antibody having human framework regions, 10 but rodent hypervariable regions. Ways of producing chimaeric antibodies are discussed for example by Morrison *et al* in PNAS, **81**, 6851-6855 (1984) and by Takeda *et al* in Nature, **314**, 452-454 (1985).

Synthetic constructs also include molecules comprising an additional moiety that provides the molecule with some desirable property in addition to antigen binding. For example the moiety may be a label (e.g. a fluorescent or radioactive label). Alternatively, it may be a pharmaceutically active agent.

Antibodies, or derivatives thereof, find use in detection/diagnosis of *S.pneumoniae*. Thus, 20 in another aspect the present invention provides a method for the detection/diagnosis of *S.pneumoniae* which comprises the step of bringing into contact a sample to be tested and antibodies capable of binding to one or more proteins described herein, or to homologues, derivatives and/or fragments thereof.

In addition, so-called "Affibodies" may be utilised. These are binding proteins selected from combinatorial libraries of an alpha-helical bacterial receptor domain (Nord *et al.*,). Thus, Small protein domains, capable of specific binding to different target proteins can be selected using combinatorial approaches.

It will also be clear that the nucleic acid sequences described herein may be used to detect/diagnose *S.pneumoniae*. Thus, in yet a further aspect, the present invention provides a method for the detection/diagnosis of *S.pneumoniae* which comprises the step of bringing into contact a sample to be tested with at least one nucleic acid sequence as described herein. Suitably, the sample is a biological sample, such as a tissue sample or a sample of blood or saliva obtained from a subject to be tested. Such samples may be pre-treated before being used in the methods of the invention. Thus, for example, a sample may be treated to extract DNA. Then, DNA probes based on the nucleic acid sequences described herein (ie usually fragments of such sequences) may be used to detect nucleic acid from *S.pneumoniae*.

In additional aspects, the present invention provides:

- (a) a method of vaccinating a subject against *S.pneumoniae* which comprises the step of administering to a subject a protein or polypeptide of the invention, or a derivative, homologue or fragment thereof, or an immunogenic composition of the invention;
- (b) a method of vaccinating a subject against *S.pneumoniae* which comprises the step of administering to a subject a nucleic acid molecule as defined herein;
- (c) a method for the prophylaxis or treatment of *S.pneumoniae* infection which comprises the step of administering to a subject a protein or polypeptide of the invention, or a derivative, homologue or fragment thereof, or an immunogenic composition of the invention;
- (d) a method for the prophylaxis or treatment of *S.pneumoniae* infection which comprises the step of administering to a subject a nucleic acid molecule as defined herein;

- (e) a kit for use in detecting/diagnosing *S.pneumoniae* infection comprising one or more proteins or polypeptides of the invention, or homologues, derivatives or fragments thereof, or an antigenic composition of the invention; and
- 5 (f) a kit for use in detecting/diagnosing *S.pneumoniae* infection comprising one or more nucleic acid molecules as defined herein.

Given that we have identified a group of important proteins, such proteins are potential targets for anti-microbial therapy. It is necessary, however, to determine whether each  
10 individual protein is essential for the organism's viability. Thus, the present invention also provides a method of determining whether a protein or polypeptide as described herein represents a potential anti-microbial target which comprises antagonising, inhibiting or otherwise interfering with the function or expression of said protein and determining whether *S.pneumoniae* is still viable.

15 A suitable method for inactivating the protein is to effect selected gene knockouts, ie prevent expression of the protein and determine whether this results in a lethal change. Suitable methods for carrying out such gene knockouts are described in Li *et al*,  
*P.N.A.S.*, **94**:13251-13256 (1997) and Kolkman *et al*, **178**:3736-3741 (1996).

20 In a final aspect the present invention provides the use of an agent capable of antagonising, inhibiting or otherwise interfering with the function or expression of a protein or polypeptide of the invention in the manufacture of a medicament for use in  
25 the treatment or prophylaxis of *S.pneumoniae* infection.

As mentioned above, we have used a bacterial expression system as a means of identifying those proteins which are surface associated, secreted or exported and thus, would find use as antigens.

The information necessary for the secretion/export of proteins has been extensively studied in bacteria. In the majority of cases, protein export requires a signal peptide to be present at the N-terminus of the precursor protein so that it becomes directed to the translocation machinery on the cytoplasmic membrane. During or after translocation, the signal peptide is removed by a membrane associated signal peptidase. Ultimately the localization of the protein (i.e. whether it be secreted, an integral membrane protein or attached to the cell wall) is determined by sequences other than the leader peptide itself.

We are specifically interested in surface located or exported proteins as these are likely to be antigens for use in vaccines, as diagnostic reagents or as targets for therapy with novel chemical entities. We have therefore developed a screening vector-system in *Lactococcus lactis* that permits genes encoding exported proteins to be identified and isolated. We provide below a representative example showing how given novel surface associated proteins from *Streptococcus pneumoniae* have been identified and characterized. The screening vector incorporates the staphylococcal nuclease gene *nuc* lacking its own export signal as a secretion reporter. Staphylococcal nuclease is a naturally secreted heat-stable, monomeric enzyme which has been efficiently expressed and secreted in a range of Gram positive bacteria (Shortle, *Gene*, **22**:181-189 (1983); Kovacevic *et al.*, *J. Bacteriol.*, **162**:521-528 (1985); Miller *et al.*, *J. Bacteriol.*, **169**:3508-3514 (1987); Liebl *et al.*, *J. Bacteriol.*, **174**:1854-1861 (1992); Le Loir *et al.*, *J. Bacteriol.*, **176**:5135-5139 (1994); Poquet *et al.*, *J. Bacteriol.*, **180**:1904-1912 (1998)).

Recently, Poquet *et al.* ((1998), *supra*) have described a screening vector incorporating the *nuc* gene lacking its own signal leader as a reporter to identify exported proteins in Gram positive bacteria, and have applied it to *L. lactis*. This vector (pFUN) contains the pAMβ1 replicon which functions in a broad host range of Gram-positive bacteria in addition to the ColE1 replicon that promotes replication

in *Escherichia coli* and certain other Gram negative bacteria. Unique cloning sites present in the vector can be used to generate transcriptional and translational fusions between cloned genomic DNA fragments and the open reading frame of the truncated nuc gene devoid of its own signal secretion leader. The *nuc* gene makes an ideal reporter gene because the secretion of nuclease can readily be detected using a simple and sensitive plate test: Recombinant colonies secreting the nuclease develop a pink halo whereas control colonies remain white (Shortle, (1983), *supra*; Le Loir *et al.*, (1994), *supra*).

Thus, the invention will now be described with reference to the following representative example, which provides details of how the proteins, polypeptides and nucleic acid sequences described herein identified as antigenic targets.

We describe herein the construction of three reporter vectors and their use in *L. lactis* to identify and isolate genomic DNA fragments from *Streptococcus pneumoniae* encoding secreted or surface associated proteins.

#### EXAMPLE 1

##### 20 (i) Construction of the pTREP1-nuc series of reporter vectors

###### (a) Construction of expression plasmid pTREP1

The pTREP1 plasmid is a high-copy number (40-80 per cell) theta-replicating gram positive plasmid, which is a derivative of the pTREX plasmid which is itself a derivative of the previously published pIL253 plasmid. pIL253 incorporates the broad Gram-positive host range replicon of pAMβ1 (Simon and Chopin, *Biochimie*, 70:559-567 (1988)) and is non-mobilisable by the *L. lactis* sex-factor. pIL253 also lacks the *tra* function which is necessary for transfer or efficient mobilisation by

conjugative parent plasmids exemplified by pIL501. The Enterococcal pAM $\beta$ 1 replicon has previously been transferred to various species including *Streptococcus*, *Lactobacillus* and *Bacillus* species as well as *Clostridium acetobutylicum*, (Oultram and Klaenhammer, *FEMS Microbiological Letters*, 27:129-134 (1985); Gibson *et al.*, 5 {FULL REF NEEDED} 1979; LeBlanc *et al.*, *Proceedings of the National Academy of Science USA*, 75:3484-3487 (1978)) indicating the potential broad host range utility. The pTREP1 plasmid represents a constitutive transcription vector.

The pTREX vector was constructed as follows. An artificial DNA fragment 10 containing a putative RNA stabilising sequence, a translation initiation region (TIR), a multiple cloning site for insertion of the target genes and a transcription terminator was created by annealing 2 complementary oligonucleotides and extending with Tfl DNA polymerase. The sense and anti-sense oligonucleotides contained the recognition sites for NheI and BamHI at their 5' ends respectively to facilitate 15 cloning. This fragment was cloned between the XbaI and BamHI sites in pUC19NT7, a derivative of pUC19 which contains the T7 expression cassette from pLET1 (Wells *et al.*, *J. Appl. Bacteriol.*, 74:629-636 (1993)) cloned between the EcoRI and HindIII sites. The resulting construct was designated pUCLEX. The 20 complete expression cassette of pUCLEX was then removed by cutting with HindIII and blunting followed by cutting with EcoRI before cloning into EcoRI and SacI (blunted) sites of pIL253 to generate the vector pTREX (Wells and Schofield, *In Current advances in metabolism, genetics and applications-NATO ASI Series, H* 98:37-62 (1996)). The putative RNA stabilising sequence and TIR are derived from the *Escherichia coli* T7 bacteriophage sequence and modified at one nucleotide 25 position to enhance the complementarity of the Shine Dalgarno (SD) motif to the ribosomal 16s RNA of *Lactococcus lactis* (Schofield *et al.*: pers. coms: University of Cambridge Dept. Pathology.)

A *Lactococcus lactis* MG1363 chromosomal DNA fragment exhibiting promoter activity which was subsequently designated P7 was cloned between the EcoRI and BglII sites present in the expression cassette, creating pTREX7. This active promoter region had been previously isolated using the promoter probe vector pSB292 (Waterfield *et al.*, *Gene*, **165**:9-15 (1995)). The promoter fragment was amplified by PCR using the Vent DNA polymerase according to the manufacturer.

The pTREP1 vector was then constructed as follows. An artificial DNA fragment which included a transcription terminator, the forward pUC sequencing primer, a promoter multiple cloning site region and a universal translation stop sequence was created by annealing two overlapping partially complementary synthetic oligonucleotides together and extending with sequenase according to manufacturers instructions. The sense and anti-sense (pTREP<sub>F</sub> and pTREP<sub>R</sub>) oligonucleotides contained the recognition sites for EcoRV and BamHI at their 5' ends respectively to facilitate cloning into pTREX7. The transcription terminator was that of the *Bacillus penicillinase* gene, which has been shown to be effective in *Lactococcus* (Jos *et al.*, *Applied and Environmental Microbiology*, **50**:540-542 (1985)). This was considered necessary as expression of target genes in the pTREX vectors was observed to be leaky and is thought to be the result of cryptic promoter activity in the origin region (Schofield *et al.* pers. coms. University of Cambridge Dept. Pathology.). The forward pUC primer sequencing was included to enable direct sequencing of cloned DNA fragments. The translation stop sequence which encodes a stop codon in 3 different frames was included to prevent translational fusions between vector genes and cloned DNA fragments. The pTREX7 vector was first digested with EcoRI and blunted using the 5' - 3' polymerase activity of T4 DNA polymerase (NEB) according to manufacturer's instructions. The EcoRI digested and blunt ended pTREX7 vector was then digested with Bgl II thus removing the P7 promoter. The artificial DNA fragment derived from the annealed synthetic oligonucleotides was

then digested with EcoRV and Bam HI and cloned into the EcoRI(blunted)-Bgl II digested pTREX7 vector to generate pTREP. A *Lactococcus lactis* MG1363 chromosomal promoter designated P1 was then cloned between the EcoRI and BglII sites present in the pTREP expression cassette forming pTREP1. This promoter was 5 also isolated using the promoter probe vector pSB292 and characterised by Waterfield *et al.*, (1995), *supra*. The P1 promoter fragment was originally amplified by PCR using vent DNA polymerase according to manufacturers instructions and cloned into the pTREX as an EcoRI-BglII DNA fragment. The EcoRI-BglII P1 promoter containing fragment was removed from pTREX1 by restriction enzyme 10 digestion and used for cloning into pTREP (Schofield *et al.* pers. coms. University of Cambridge, Dept. Pathology.).

(b) **PCR amplification of the *S. aureus* nuc gene.**

15 The nucleotide sequence of the *S. aureus* nuc gene (EMBL database accession number V01281) was used to design synthetic oligonucleotide primers for PCR amplification. The primers were designed to amplify the mature form of the nuc gene designated nucA which is generated by proteolytic cleavage of the N-terminal 19 to 21 amino acids of the secreted propeptide designated Snase B (Shortle, (1983), 20 *supra*). Three sense primers (nucS1, nucS2 and nucS3, Appendix 1) were designed, each one having a blunt-ended restriction endonuclease cleavage site for EcoRV or SmaI in a different reading frame with respect to the nuc gene. Additionally BglII and BamHI were incorporated at the 5' ends of the sense and anti-sense primers respectively to facilitate cloning into BamHI and BglII cut pTREP1. The sequences 25 of all the primers are given in Appendix 1. Three nuc gene DNA fragments encoding the mature form of the nuclease gene (NucA) were amplified by PCR using each of the sense primers combined with the anti-sense primer described above. The nuc gene fragments were amplified by PCR using *S. aureus* genomic DNA template,

Vent DNA Polymerase (NEB) and the conditions recommended by the manufacturer.

An initial denaturation step at 93 °C for 2 min was followed by 30 cycles of denaturation at 93 °C for 45 sec, annealing at 50 °C for 45 seconds, and extension at 73 °C for 1 minute and then a final 5 min extension step at 73 °C. The PCR amplified products were purified using a Wizard clean up column (Promega) to remove unincorporated nucleotides and primers.

5                   (c) Construction of the pTREP1-nuc vectors

10          The purified nuc gene fragments described in section b were digested with Bgl II and BamHI using standard conditions and ligated to BamHI and BglII cut and dephosphorylated pTREP1 to generate the pTREP1-nuc1, pTREP1-nuc2 and pTREP1-nuc3 series of reporter vectors. General molecular biology techniques were carried out using the reagents and buffer supplied by the manufacturer or using 15 standard conditions(Sambrook and Maniatis, (1989), *supra*). In each of the pTREP1-nuc vectors the expression cassette comprises a transcription terminator, lactococcal promoter P1, unique cloning sites (BglII, EcoRV or SmaI) followed by the mature form of the nuc gene and a second transcription terminator. Note that the sequences required for translation and secretion of the nuc gene were deliberately excluded in 20 this construction. Such elements can only be provided by appropriately digested foreign DNA fragments (representing the target bacterium) which can be cloned into the unique restriction sites present immediately upstream of the *nuc* gene.

In possessing a promoter, the pTREP1-nuc vectors differ from the pFUN vector described by Poquet *et al.* (1998), *supra*, which was used to identify *L. lactis* exported proteins by screening directly for Nuc activity directly in *L. lactis*. As the pFUN vector does not contain a promoter upstream of the *nuc* open reading frame the cloned genomic DNA fragment must also provide the signals for transcription in

addition to those elements required for translation initiation and secretion of Nuc. This limitation may prevent the isolation of genes that are distant from a promoter for example genes which are within polycistronic operons. Additionally there can be no guarantee that promoters derived from other species of bacteria will be recognised and functional in *L. lactis*. Certain promoters may be under stringent regulation in the natural host but not in *L. lactis*. In contrast, the presence of the P1 promoter in the pTREP1-nuc series of vectors ensures that promoterless DNA fragments (or DNA fragments containing promoter sequences not active in *L. lactis*) will still be transcribed.

10

**(d) Screening for secreted proteins in *S. pneumoniae***

Genomic DNA isolated from *S. pneumoniae* was digested with the restriction enzyme Tru9I. This enzyme which recognises the sequence 5'- TTAA -3' was used because it cuts A/T rich genomes efficiently and can generate random genomic DNA fragments within the preferred size range (usually averaging 0.5 - 1.0 kb). This size range was preferred because there is an increased probability that the P1 promoter can be utilised to transcribe a novel gene sequence. However, the P1 promoter may not be necessary in all cases as it is possible that many Streptococcal promoters are recognised in *L. lactis*. DNA fragments of different size ranges were purified from partial Tru9I digests of *S. pneumoniae* genomic DNA. As the Tru 9I restriction enzyme generates staggered ends the DNA fragments had to be made blunt ended before ligation to the EcoRV or SmaI cut pTREP1-nuc vectors. This was achieved by the partial fill-in enzyme reaction using the 5'-3' polymerase activity of Klenow enzyme. Briefly Tru9I digested DNA was dissolved in a solution (usually between 10-20 µl in total) supplemented with T4 DNA ligase buffer (New England Biolabs; NEB) (1X) and 33 µM of each of the required dNTPs, in this case dATP and dTTP. Klenow enzyme was added (1 unit Klenow enzyme (NEB) per µg of

DNA) and the reaction incubated at 25°C for 15 minutes. The reaction was stopped by incubating the mix at 75°C for 20 minutes. EcoRV or SmaI digested pTREP-nuc plasmid DNA was then added (usually between 200-400 ng). The mix was then supplemented with 400 units of T4 DNA ligase (NEB) and T4 DNA ligase buffer (1X) and incubated overnight at 16°C. The ligation mix was precipitated directly in 100% Ethanol and 1/10 volume of 3M sodium acetate (pH 5.2) and used to transform *L. lactis* MG1363 (Gasson, 1983). Alternatively, the gene cloning site of the pTREP-nuc vectors also contains a BglII site which can be used to clone for example Sau3AI digested genomic DNA fragments.

10 *L. lactis* transformant colonies were grown on brain heart infusion agar and nuclease secreting (Nuc<sup>+</sup>) clones were detected by a toluidine blue-DNA-agar overlay (0.05 M Tris pH 9.0, 10 g of agar per litre, 10 g of NaCl per liter, 0.1 mM CaCl<sub>2</sub>, 0.03 % wt/vol. salmon sperm DNA and 90 mg of Toluidine blue O dye) essentially as described by Shortle, 1983, *supra* and Le Loir *et al.*, 1994, *supra*). The plates were 15 then incubated at 37°C for up to 2 hours. Nuclease secreting clones develop an easily identifiable pink halo. Plasmid DNA was isolated from Nuc<sup>+</sup> recombinant *L. lactis* clones and DNA inserts were sequenced on one strand using the NucSeq sequencing primer described in Appendix 1, which sequences directly through the DNA insert.

20 **Isolation of Genes Encoding Exported Proteins from  
*S. pneumoniae***

A large number of gene sequences putatively encoding exported proteins in *S. pneumoniae* have been identified using the nuclease screening system. These have 25 now been further analysed to remove artefacts. The sequences identified using the screening system have been analysed using a number of parameters:

1. All putative surface proteins were analysed for leader/signal peptide sequences using the software programs Sequencher (Gene Codes Corporation) and DNA Strider (Marck, *Nucleic Acids Res.*, 16:1829-1836 (1988)). Bacterial signal peptide sequences share a common design. They are characterised by a short 5 positively charged N-terminus (N region) immediately preceding a stretch of hydrophobic residues (central portion-h region) followed by a more polar C-terminal portion which contains the cleavage site (c-region). Computer software is available which allows hydropathy profiling of putative proteins and which can readily identify the very distinctive hydrophobic portion (h-region) typical of leader peptide 10 sequences. In addition, the sequences were checked for the presence of or absence of a potential ribosomal binding site (Shine-Dalgarno motif) required for translation initiation of the putative nuc reporter fusion protein.

2. All putative surface protein sequences were also matched with all of the protein/DNA sequences using the publicly databases [OWL-proteins inclusive of 15 SwissProt and GenBank translations]. This allows us to identify sequences similar to known genes or homologues of genes for which some function has been ascribed. Hence it has been possible to predict a function for some of the genes identified using the LEEP system and to unequivocally establish that the system can be used to 20 identify and isolate gene sequences of surface associated proteins. We should also be able to confirm that these proteins are indeed surface related and not artifacts. The LEEP system has been used to identify novel gene targets for vaccine and therapy.

3. Some of the genes identified proteins did not possess a typical leader peptide sequence and did not show homology with any DNA/protein sequences in the database. Indeed these proteins may indicate the primary advantage of our 25 screening method, i.e. the isolation of atypical surface-related proteins, which may have been missed in all previously described screening protocols or approaches based on sequence homology searches.

In all cases, only partial gene sequences were initially obtained. Full length genes were obtained in all cases by reference to the TIGR *S.pneumoniae* database ([www.tigr.org](http://www.tigr.org)). Thus, by matching the originally obtained partial sequences with the database, we were able to identify the full length gene sequences. In this way, as described herein, three groups of genes were clearly identified, ie a group of genes encoding previously unidentified *S.pneumoniae* proteins, a second group exhibiting some homology with known proteins from a variety of sources and a third group which encoded known *S.pneumoniae* proteins, which were, however, not known as antigens.

## Appendix I - Oligonucleotide primers

nucS1

Bgl II Eco RV

5' - cgagatctgatatctcacaaacagataacggcgtaaatag -3'

nucS2

Bgl II Sma I

5' - gaagatctccccggatcacaaacagataacggcgtaaatag -3'

10

nucS3

Bgl II Eco RV

5'- cgagatctgatatccatcacaacagataacggcgtaaatag -3'

15

nucR

## Bam HI

5' - cggatcctatggacctgaatcagcggtgtc -3'

NucSeq

5'- ggatgc~~t~~ttgtttcagggtgtatc -3'

pTREPF

5' - catgatatcggtacctaagctatatcattgtccggcaatggtggtggcttttttgttttagcggataa  
caatttcacac -3'

25

pTREPR

5'- gcggatccccgggcttaattaaacactagtcgaagatctcgcaattctccgtgtgaaatt  
gttatccgcta -3'

30

pUCF

5' - cgccagggtttcccagtcacgac -3'

VR

5'- tcagggggcgaggacctatg -3'

35

V1

5' - tcgtatgttgtgtggaaatttg -3'

V2

5'- tccggctcgatgttgtgaaattg -3'

TABLE 1

ID4 1200 bp

5 ATGAGAAAATATGTGGTTGTAATCAAGGAAACCTATCTTCGACATGTCGAGTCATGGAGTTCTTCTTTATGGTGA  
 TTTCGCCGTTCCCTCTTTAGGAATCTCTGTAGGAATTGGCATCTCCAAGGTTCTCTATGGCTAAAAATAATAAAA  
 GTGGCAGTAGTGACAACAGTGCCATCTGTAGCAGAAGGACTGAAGAATGTAATGGTAACTTCGACTATAAA  
 GACGAAGCAAGTGCCTAACAGCAATTAAAGAAGAAAAATTAAAGGTTATTGACCATTGATCAAGAAGATAGT  
 GTTCTAAAGGCAGTTATCATGGCAAACATCGCTGAAAATGGAATTAAATTGAGGTTACAGGTACACTCAATG  
 AACTGCAAACATCAGCTTAATCGTCAACTGCTCCTGTCTCAAGAGCAGGAAAACGCTTAGCGCAGACAATTCA  
 ATTACAGAAAAGATTGATGAAGCAAGGAAAATAAAAGTTTCTCAAACAATTGCAAGCAGGTGCCTTAGGATT  
 CTTCCTTATGATTCTGATTACCTATGCGGTGTAACAGCTCAGGAAGTGCAGGCTGAAAAGGCACCAAAATT  
 10 ATGGAAGTCGTTCTTCTAGCATAAGGGCAAGTCACTATTCTATGCGGGATGATGGCTCTGTTCTAGTGTGTT  
 AACGCATATTGGGATCTATGTTGAGGTGGCTGGCTGGCTTCTGCTCTTAAAGATTGCACTTCTGGCTCAGT  
 15 CTGGTATTGGATCACCTGGGAGATGCTATCTCACTGAATACCTTGCTCTTATTGATCAGTCTTCTATGTAC  
 GTAGTCCTGGCAGCCTCTAGGATCTATGGTTCTGCTCTGAGGACTCAGGGAAAGCCTGTCGCCTTGATGA  
 TTTGATTATGGGTGGTTTTGGAGTGACAGCTCTAGGTGCAGCTGGTGACAATCTCCTTGAAGATTGGTTCT  
 20 TATACTCCCTTATTGACCTCTTATGCCGTTCTGAACGATTAATGACTATGCGGGGGGAGCAGAAGCATGGA  
 TTTCACTTGCTATTACAGTGATTTGCGGTGTTAGCAACAGGATTATCGGACGCATGTATGCTAGTCGTTCT  
 CAAACGGATGATTAGGGATTGGAAAACCTTAAACGTCCTTATCTTAAAG  
 MRNMWVVIKETYLRHVESWSFFFMVISPLFLGLVISVGIGHLQGSSMAKNKVAVVITVPSVAEGLKNVNGVNFDYKDE  
 25 ASAKEAIKEEKLKGYLTDQEDSVLKAVYHGETSENGIKFETGTLNELQNLNRSTASLSQEKEKLAQTIQFTEKIDE  
 AKENKKFIQTIAAGALGFFLYMILITYAGVTAQEVASEKGTKIMEVFSSIRASHFYARMMALFLVILTHIGIYVVG  
 AAVLLFKDLPFLAQSGILDHLGDAISLNLLFILISLFMFYVLAFLGSMVSRPEDSGKALSPMLIMGGFFGVTLGAA  
 GDNLLLKIGSYIPFISTFFMPFRINDYAGGAEAWISLAITVIFAVVATGFIGRMYASLVLTQDDLGWKTFRKRALSYKZ

ID5 1125 bp

30 CCTGGAAAGTCCTGAAAATTATGATAGAATGGTGAAGGAAAATTCAAGGAGAGTAGTAGTGACTCAAATGTT  
 GAAAGTCCTCGTATCCATTGTAATCAGTCATACAATGAAGAAAATATCTGCCCTGGTCTAATTGAAGACTAA  
 AAAATCAAACCTATCTAAAGAGGATATTGAAATTCTTAAATGCTATGTCACAGGGACACAGCTAT  
 35 CATTCACTTAAAGGAGATACAGAGTTAACTCAATTAGATGTATAACATCTAAAGAAAATCAAGC  
 TAGTGGTTTAACCTGGGAGTTAACATTCTGTAGGGACCTTATTAAAATTGATGCTCATTCAAAGTTACT  
 GAGACTTTGTAATGAAACATTGTCATTATTCAACAAGGTGAATTGCTGTGGGGGCTAGACCGACGATTG  
 TCGAAGGAAAAGGAAATGGGAGACCTGTCATCTGTTGAGGAAAATATGTTGGCAGTAGCATTGCCAATT  
 ATCGAAATAGTCTGAGGGATAGATATGTTCTCTATTTCATGGAATGTATAACGAGAGGTTTCCAGAAGGT  
 40 TGGTTTAGTAAATGAGCAACTTGGCGAAGTGAAGATAATGATAATTGATTAGAATTGAGAATATGGTTATAAA  
 ATCCGCTATAGCCCAAGTATTCTATCAGTATATTGACCAACATTCAAGAAAATGCTGATCAAAGTATT  
 CAAATGGTTGTGGATTGGCTTGACAAGTCATGTCAGTTAAGTGTGTTATCATTATTCACTATGTTCTGTTA  
 TTTGTTTGTGATTGTGTTAGTCAGTATTGACATTGTCACATTGTTACTATGTTACTATTAGGTGCTTATT  
 45 TTCTACTTTGTCATTACTCACTTGTGACTTTATTAAACATAAAAATGGTTCTAATTGATGCCCTTATT  
 TTATTTCACCTCACTTGCTATTGGCTTGGGACATTGAGGTTAATTAGAGGATTAAATGGAAGAAGGAGT  
 ACAAGAGAACATAATTATTGGATAAAATAAGCCAATAAAATGCTATAA  
 PGKVLKIMIEWKEKFRVVVTQNVESLLSIVISAYNEEKYLPGLIEDLKNQTPKEDIEILFINAMSTDGTTAIQQFIK  
 EDTEFNSIRLYNNPKKNQASGFNLGVKHSGVDLILKIDAHSKVTFVMNNVAIIQGEFVCGGPRPTIVEGKGKWAET  
 LHLVEENMGSSIANRNSSEDRYVSSIFHGMYKREVFKVGLVNEQLGRTEDNDIHYRIREYGYKIRYSPSILSYQYIRP  
 50 TFKKMLHQKYSNGLWIGLTSHVQFKCLSLFHYPCLFVLSLVSALLPITFVFTLLGAYFLLSLLTLLKHKNGF  
 LVMPFILFSIHFAYGLGTIVGLRGFKWKKEYKRTIYLDKISQINQNMIZ

ID11 696 bp

55 ATCATGAAAGAACAAAATACGATAGAATCGATGTTTCAATTAGTTAAAGCTTGTGAAACGCAAGCTAATG  
 ATTAAATAGTGGCACTTGTGACAGGTGCGGGGGCTTTGCAATAGCAGTTATTGTTAAAGCCAGAATATACGA  
 GTACCACCGAATTACGTAGTGAATCGCAATCAAGGAGACAAGCCGGGGTGCACAAATCAGGATTGCAAGGCAG  
 GAACCTTATCTGGTAAAGACTACCGTGAGATTATCCTTCGCAAGGATGTTGGAGGAAGTTGTTCTGATTGAA  
 ACTAGATTGACGCCAAGGTTGGCTAATAAAATTAAAGTGACAGTACCACTGTTGAGAGAAGTACGTTGCTAATT  
 TCAGTTAATGATCGAGITCCAGAAGAGGCAAGCCGTATCGCTAACTCTTGAGAGAAGTACGTTGCTCAA  
 60 AAAATTACCTGTTGACGTCACACTGGAGGAGGCAAGGCCGGCATATCCCCGCTTGCACAAATAT  
 TAAACGCAAAACACTAATTGGTTTTGGCAGGGGTGATTGAAACTAGTGTATAGTTCITCATCTTGAACTTTGG

ATACTCGTGTGAAACGTCCGGAAAGATATCGAAAATACATTGCAGATGACACTTTGGGAGTTGTGCCAAACTTGG  
GTAAGTTGAAATAG

MMKEQNTIEIDVFQLVKSLWKRKLMILIVALVTGAGAFAYSTFIVKPEYSTRYVVNRNQGDKPGLTNQDLQAGTYL  
VKDYREIILSQDVLEEVVSDLKLDDTPKGLANKIKVTVPDTRVISVNDRVEEASRIANSLREVAAQKIIISITRSDVTT  
LEEARPAISPSSPKRNLIGFLAGVIGTSVIVLHLELLDTRVRPEDIENTLQMPLLGVVPNLGKLKZ

ID19 555 bp

10 ATGGTAAAAGTAGCAGTTATATTAGCTCAGGGCTTGAAGAAAATTGAAGCCTTGACAGTTGAGATGTCCTGCGTC  
GAGCCAATATCACATGTGATATGGTTGGTTGAAGAGCAAGTAACGGGTCGATGCAATCCAAGTAAGAGCAG  
ATCATGTCTTGATGGAGATTATCAGACTATGATATGATTTCTTCCTGGAGGTATGCCTGGTTCTGCACATTTA  
CGTATAATCAGACCTGATTCAAGAATTGCAAAGCTTCGAGCAAGAAGGGAAAGAAACTAGCAGCCATTGTCG  
GCACCAATTGCCCTCAATCAAGCAGAGATATTGAAAATAAGCATACTTGTATGACGGCGTTCAAGAGCAA  
15 ATCCTGATGGTCACTACGTCAAGGAAACAGTAGTGGTAGATGGTCAAGTGCACACCAGTCGGGTCTCAACA  
GCCCTGCTTGCCTACGGAGTTGGTAGGCAACTAGGAGGGACGCAGAGAGTTACGAACAGGAATGCTTAT  
CGAGATGCTTGGTAAAATCAGTAA

20 MVKVAVILAQQFEEIEALTVVDVLRRANITCDMVGEEQVTGSHAIQVRADHVFDGDLSDYDMIVLPGGMPGSAHLR  
DNQTLIQELQSFEQEGKKLAAICAAPIALNQAEILKNKRYTCYDGVQEILDGHYVKETVVVDGQLTTSRGPSATALAFA  
YELVEQLGGDAESLRGMLYRDVFGKNQZ

ID27 306 bp

25 GTGGTAGGGATGGTAGAACCAACCTAGAAAGCCTTATAAAAGATCTTACAATCATGCTGACATGATTGAGT  
GAAGATTAGTTGCTGCTCTCTAGAGACTACTAAAAACTGCCTACTACAAATGAGCAATTGCAAGGAGTCGTC  
TCTCAGGCCTGGTCAATCGTAATTGCTCTAAATCCCAACATCCAGCACCTGAGTTGCTCAACTGGCTGCTT  
TGTCAAAAGAGAAGAACGAAAGTACAGAGGAACACTGCACCTCTCGCCTATGATGAGGAACCTTTAAATGCT  
TTGA

30 MVGMVEPNLESLIKDLYNHARHDLSEDLVAALETTKLPTTNEQLQAVRLSGLVNRELLNPKHAPELLNLARFK  
REEAKYRGTATSALMYEELFKMLZ

ID29 945 bp

35 TTGTTCTTAAAAAGAACGAGAGGTAATCAGCATGCGTAAATGGACAAAAGGATTCTCATCTTGGGTGGTG  
ACTACCGTTATCGGTTTATCCTGCTTTTGATGGTATCCAATCTGACGGGAATAAGACCTACTTCCATGTC  
AGAACCTGCTATGATAGCCGTACGGAAAGCTAACCTTGGCAAGGAAGTCGAAAACCTAGAAATTACTCTCA  
CCAACACACCGTCAACATCACAGACTCTTCGATGATCAAATCCACATTCTTACCATCTCTCTGCTCACC  
40 ATGATCTTATCACAATCAGAACGATAGAAACTGAGTCACTGATAAGAAACTGCTGAAACTCCGTTCTC  
TCTGGAATTGGGGATTCTCATATCGCAAGTAGCTACTCTAGTCGTTTGAGAAGTTATCTCCGACTAAC  
AAAGGGAGAACTCTAAAGGGATCAACATCTCAGCAATCGCGGACAACCACCATCATAATGCTAGCCTTGAA  
AATGCGACCCCAATACAAACAGCTATATCTCCGAATTGAAGGAAGTCGATACAAAACAGTAAACTCACACG  
CCAATATGTTAATATCTTGTACAGCTTACAGATAGTCAGTAGCTCAAGGAAATCCTCCACGCTG  
45 AAAATATCCAAGTCCATGGCAAGGTTGACTGCTTCAAGAACATGGTTATCTCAGAATCATCTCTGAC  
GCCAACGAAATTACTGGACATCTCAAGAACATGGTTATCTCAGAATCATCTCTGAC  
GAGGTACGGAAATTAGCAACCCCTAACAAACTGAAAGAACCGATGTCAAGGATCAACTCATTCGAGATCTGATG  
ATAATATTGATCTAATATCACACCAAGCAGACGTTGA

50 MFLKKEREVISMRKWTKGFLIFGVTTVIGFILLFVGIQSDGIKSLLSMSKEPVYDSRTEKLTFGKEVENLEITLHQHTLT  
TDSFDDQIHISYHPSLSAHDLITQNDRTLSLTDKLLSETPLSSGIGGILHIASSYSSRFEEVILRPLPKGRTLKGINISANR  
GOTTINASLENATLNTNSYLRIEGSRIKNSKLTTPNIVNIFDTLTDSQLESTENHFHAENIQVHGKVELTAKDYLRIID  
QKESQRINWDISSNYGSIFQFTREKPESRGTELSNPYKTEKTDVKDQLIARSDDNDLISTPSRRZ

ID30 879 bp

55 ATGAAACAAAGAATGGTTGAAGTAATGATTGTAAAGAACACAAGCAAGAACAGCTGAAGAGCAAGCTCAA  
GAGGTTGCAGACAAGGCTGAAGAACAGATAGCGATCTCGATACACCAATTGAAAAGAAATACTEAGTTAGAGGAG  
GAAGTCCCCTCAAGCTGAAGTCGAATTGGAAAGGCCAGCAAGAACAGAAAATTGAAGCTCTGAAAGACAGTGAAGC  
GAGAACAGAAATTAGAACAAAGGCAAGCTAACATTCTACTGAAGAACAGGCCACTCTTCTAAAGAACAGAAA  
60 AAGTCACATAGCTGAAGAGAGCAGAACAGCTCTCTCAGCAAAAAGCAACCACGAAAGAGCCACTCTTATCA  
GTAATTCTTATGAAAGCTTATATCCCCGACCAAGCTCAAATCTAGGGATAATGGAAAGAGCAAGTGCCTG  
ATTTCGGCTCTGGCTAGTGGAAAGCGATCAAATCTCTACAAGTAAGTGGAAACAAGTACACACACAGTTACAC  
ACCCCTCTCTGCTATTCTGTTCTGATCTCTCTTCTTCTAGTATCTATCACATCAAACATGCTTACTATGG

ACATATAGCAAGCATTAAACAGTCGCTTCCCTGAGCAGCTAGCTCCTTAACCTTTCTATCATCTATCCTAG  
 TAGCGACAACACTCTTCTCTTTCATCCCTTGGGTAGTTCTGAGACGATITATCCACCAGGAAAAGGA  
 CTGGACGCTAGACAAGGTTCTCAACAATATACTGCAACTCTGGCAATCCAACTCTCCTACTGCTATTGCTAGTT  
 TCTTGCCTTGTGATAGCCTACGATTACAGCCCCTTGTGTGA

5 MKQEWFESNDFVKTTSKNPKPEEQAEVADKAEEIADLDPIEKNTQLEEVPAEVELESQEEKIEAPEDSEARTEIE  
 EKASNSTEEPDLSKETEKVTIAEESQEALPQQKATTKEPLLISLESPYIPDQAPKSRDKWKEQVLDFWSWLVEAIKS  
 PTSKLETTSITHSYTAFLLLILFSASSFFSIYHIKHYGHIASINSRFPQLAPLTLFSIILVATTLFFFLLGSFVRRFIH  
 QEKDWTLDKVLLQQYSQLAIPISSLLLVSLLSIIAYDLQPSCVZ

10 **ID105 990 bp**  
 ATGCAACTCGCTTCTCGGTCTACTCATTGTCGCTGGTACAATTGTCCTAAAAAAGGAAAGAGAGGTAATCA  
 GCATGCGTAAATGGACAAAAGGATTCTCATCTTGGTGGTACTACCGTTATCGGCTTATCCTGCTTTGTA  
 GGTATCCAATCTGACGGGATTAAGAGCCTACTTCCATGTCACAGAACCTGCTATGATAGCCGTACGGAAAAG  
 CTAACCTTGGCAAGGAAGTCGAAAACCTAGAAATTACTCTCCACCAACACGCTCACCATCACAGACTTTCG  
 ATGATCAAATCCACATTCTTACCATCCATCTTCTGTCACCATGATCTTATCCAATCAGAACGATAGAAC  
 TCTGAGCTCTACTGATAAGAAACTGCTGAAACTCCGTTCTCTCTGGAATTGGTGGGATTCTCATATCGAA  
 GTAGCTACTCTAGCTGTTGAGAAGTATTCTCGACTACAAAAGGGAGAACTCTAAAGGGATCAACATCTC  
 AGGCAATGGAAGGAAGTCGTATCAAAAACAGTAAACACTCACAACGCCAATATGTTAATATCTTGATACAGTCTT  
 ACAGATAGTCAGCTAGAGTCACAGAGAATCACTCCACGCTGAAAATATCCAAGTCCATGGCAAGGTTGAAC  
 ACTGCCAAAGATTATCTCAGAATCATCTAGACAGAAAGCCAACGAATTAACTGGGACATCTCAAGCAAC  
 TATGGTCTATCTCCAATTCAACAGAGAAAAGCCTGAATCAAGAGGTACGGAAATTAAAGCAACCTTACAAA  
 GAAAAAACCGATGTCAAGGATCAACTCATTGCGAGATCTGATGATAATTGATCTAATATCCACACCAAGCAGA  
 CGTTGA

20 MQLASSVSYSLVWYNLFKKEREVISMRKWTKGLIFGVVTVIGFILLFGIQSDGIKSSLMSKEPVYDSRTEKLTFGK  
 EVENLEITLHQHTLTITDSFDDQIHISYHPSLSAHDLITQNDRTLSLTDKLSSETPFLSSGIGGILHASSYSSRFEEVILR  
 LPKGRTLKGINISANRGQTIIINASLENATLNTNSYILRIEGSRKNSKLTPNIVNIFDTVLTDQSLESTENHFHAENIQVH  
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25 **ID107 -78bp**

30 35 ATGATATGAAAATGAAGCAGGGAGGGAGCAGGGCGTGTGGGATGGAGAGTGGGGAGGGACGCTGCTATT  
 AATC

40 MICKMKQGGSRACWGWRVGRCYFN

45 **ID109 714 bp**

45 CGATAAAGAGGCCCTGAGTAATCTAATTGAGATTGAAATGGAGAGATTATGGGCTTGATTGGTCATAATGG  
 GGCTGAAAATCGACCACTATAAAATCCCTAGTCAGTATCATTCACCCAGCAGGGCGTATTGGTAGACGGT  
 CAGGAGTTATCGGAAAATCGCTGGCTATTAAACGAAAGATTGGCTACGTAGCAGACTGCCGTACTTATTTAC  
 GCTTAACGCCAATGAATTGGGATTGATCGCCTCATCTATGATCTGAGTAGCTGACTTGGAGGCTAGTCT  
 AGCTAGGCTATTGAACGTTTTGATTGCTGAAAATCGCTACGGTACGTTGAGAAGTAACTCTTCTCACGGAATGCGTC  
 AGAAAAGTCTTGTATCGGAGCAGCTCTGTGATCCCAGATATTGGGTTGGACGAACCCCTGACTGGTTGG  
 50 TCCCCAGGTGCTTGTGATTGAAACAGATGATGAAGGAACATGCACAAAAGGGAGACAGTCTTGTGTTCAAC  
 TCATGCTCTAGAGGTGGAGAGCAAGTCTGTGATCGGATTGCCATTGAAAGGGGATTGATTATTGTGGT  
 AAGGTAGAGGACTTGAGGAAAGACCACCCAGACCAGTCTTGGAAAGTATCACCTAGTCTGCTGGTAGAAAA  
 GAGGAGGTTGCGGATGCGTCAAGGTATTAA

55 DKEALSNLNQIENGIMGLIGHNGAGKSTTIKSLVSIISPSSGRILVDGQELSENRLAKRKIGYVADSPDLFLRLTANE  
 WELIASSYDLSRSDEASLARLLNVDFEAENRYQVIETLSHMRQKVFIGALLSDPDIVWLDEPLTGLDPQAADF  
 MMKEHAQKGKTVLFSTHVLEVAEQVCDRIAILKKHLYCGKVEDLRKDHPDQSLEIYLSLAGRKEEVADASQGHZ

60 **ID112 360 bp**

60 ATGGCTTGTTCAGAGAGAGGAGCAGTACGGAGACACCAATGGCAAGTCCAATAATGAGACCTATGATGGTT  
 CCGACGATAGAGATTAAAGAGTGAATCCAGCACCACGCAAGAGTTGTCAGTTGCAAGTAAAGAATTAGCA  
 ACTTGGCTAAAGAAACTACTGCTAGTCTCTCAGTTGTTAGCTTCGGCAGGTTGTTGATCATACGATCCAT

CAAGGCAACTGGTCATCTTTGAATGGTTCAATGCTGGCATTGATTGGCTAATACGATTGTCACTTACGAA  
GCCCGATAGCGATAGCTGTATCTTCTCCCAGTTGAAACCAGGTTCTACTTGA

5 MALFSERGAVRKTPMASPIMRPMMVPTIEIKRVIPAPRKSCQFSERILATWLKKLLLSSVVVASAGCSLIIRSIKATWSS  
FEMVSQLALIWLIRLSFLRSPIAIAVSSSPVLKPGSTZ

TABLE 2

ID2 840 bp

5 ATGGGAATTGCTCTAGAAAATGTGAATTACATATCAAGAAGGTACTCCCTAGCTTCAGCAGCTTGTGGATG  
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 ACTCTTAAATGGTTATTGGTGCCTAAGGAGTGTGGGTTTGTACCTTAATCACCTGACTCTAAA  
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 GAAACTGGCTCTGGTGGAAATTGATGAATCACTTTGATCGTAGTCCGTTGAGCTGTCAAGGGGGACAAAATGAGA  
 10 CGTGTGCCATTGCAGGCATACTTGCCTGGAGCCAGCTATATTAGTCTTAGATGAGCCAACAGCTGGTCAAGATC  
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 15 AACCAAGTGTGCTTCAAGACGTTGTTTATGAGAAGATTCAGTTGGGAGTACCTAAAATTACGGCCTTGT  
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MGIALEVNFTYQEGTPLASAALSDVSLIEDGSYTALIGHTGSGKSTILQLLNGLLPSQGSVRVFDTLITSTSKNKDIR  
 20 QIRKQVGLVFQFAENQIFEETVLKDVAFGPQNFGVSEEDAVKTAKEKLALVGIDESLFDRSPFELSGQMRRVIAIGILA  
 MEPAILVLDEPTAGLDPLGRKELMTLFKKLHQSGMTIVLVTHLMDDVAEYANQVYMEKRLVKGGKPSDVFQDV  
 FMEEVQLGVPKITAFCKRЛАDRGVFSKRLPKIEEFKESLNGZ

ID 3 6360 bp

25 TACCCGGTAGTCTAGCAGACACATCTAGCTCTGAAGATGCTTAAACATCTGTATAAAGAAAAAGTAGCAGAA  
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 GAAGAAAATTCAAATAATCCAAAGGAGATTACGGACTCATTTGTGAATAAAACACAGAAAATCCAAAAAA  
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 30 CTTAAGAATACAAAAGTTTATATACTTGTAGATAATTAAACGGTAGTGCCTAGAGAAAACAACACTCCAGATAACT  
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 35 TGCCAAAGCCTCAATGAGATTTAAAAGAACGACTAAAAGGCACTGATAAAAATTATTGGTTGAGTGAATAAAAT  
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 45 TTGATAGGTTGGATCTAGGGCAAATTGCACTGAGTGTAGAATTATAACAGGATTTAAAATGCTTAA  
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 50 GTDKNYLSDKIPHAFNYNGGKITVEKYDDGRDYFDPHMGHLAGILAGNDTEQDIKNFNGIDGIAPNAQIFSYKMS  
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 60 APESKDNQDPATKIRGQFEPIAEGQYFYKFYRLKDYWPWQSYIPVKIDNTAPKIVSDFSNPEKIKLITKDTYHKVKD  
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EIKSKIYGVLSPSKDGHFEILGKISNVSKNAKVYYGNNYSIEIKATKYDFHSKTMTFDLYANINDIVDGLAFAGDMRLF  
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 5 MNDKGEAIDKDGNLVTDSSKLVLFGKDDKEYTGEDKFVNVEAIKEDGSMLFIDTKPVNLSDMKNYFNPSSNKIYVRNP  
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 VYLQTGSDLNNAKAVGVHYQFLYDNVKPEVNIDPKGNTSIEYADGKSVVFNIINDKRNNGFGEIQBHQIYINGKEYTSF  
 NDIKQIIDKTLNIKIVVVKDFARNTTVKEFILNKTGEVSELKPHRTVTIQNCKEMSSTIVSEEDFILPVYKGELEKGYQFD  
 GWEISGFEGKKDAGYVINLSKDTPIKPVFKKEEENKGTFDVSKKDDNPQVNHSQLNEHRKEDLQPEEHSQKSDS  
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ID6 597 bp

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CTTGAATTAAATAAAAAACGTATGCGACTAAGCATTACTGATAAGCTTGTGATCCCAAAGATGTGCGTACGG  
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ID7 1401 bp

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MTAIDFTAEEVKEKRKEDLLADLFSLEINSERDDSKADAQHPFGPGPVKALEKFLEIADRDGYPTKNVDNYAGHFEGDG  
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ID8 1617 bp

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MYTIKSNKKFSLLTIFIVAGQLLIYAAATINALVNLNIELAMNLERFLKLSIYQMIWCGIIFLDWVVKNYQVEVIQEFNL  
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 20 LLVPKIFASKMREVSLNLTQNQEAFLKSSSETILNGFDVLASLNLLYVLPKKIKEAGILLKVMVQRKTTVETLAGAISFFLNI  
 FFQISLVFLTGYLAIKGIVKIGTIEAIGALTGVIFTALGELGGQLSSIIGTKPFLKLYSINPIESNKMNDIEPNEVNRFDFPLYE  
 AKNICYKYGDKEILKNLNFCFQRNEKYLILGESGSGKSTLLKLNGFLRDYSGELEFCGDDIKKTSYLNMVSNVLYVDQ  
 KAYLFEGTIRDNLILLEENYTDEEILQSLEQVGLSVKDFPNNILYYVGDDGRLLSGGQKQKITLARGLIRNKKIVLIDEQT  
 SAIDRRRTSLAIERKILDREDLTVIIVTHAPHPELKQYFTKIYQFPKDFIZ

**ID9 705 bp**

25 ATAACAGTTAACAGATTATGGACAAATAGCCGTTAGATATGACTGCAAGGCCTATTACAGGAATTAGCT  
 GATAAAGATTGCTGATTGCTGATGGTGGAGCTGAAAACCTTCGAACCAACTCCCTTTGACTAATGAGCGAT  
 CAAATATTGAAAAACAAGCCCTCCAAACGGCAGAAAACAAGAAAATAGCCCATTGCGAGCTAGTAAAG  
 30 AAAGAGAAAATCTTCAATTGGACCAGGAACACATTAGAGTTTGCCTGAGTTGCCTATTGACAATATCCG  
 CGCTGTAACCAACAGTCTACCTGTTCTGATTAAAGCAACGAAAATTAAACAGATTGATTAAATAGGTGGA  
 ATTATCGCGATATTACAGGTGTTCTGTTGATCATTGACCTACAAAATCTCTAATCTCAATTCTAAAGC  
 TTGTTAGCTGTAATGGTATTCAAACAGGAGCTCTAGCTACTTTAGCGAGGAAGAGGGAGAGGCTAACGCATC  
 GCTTAAATAATTCTAATAAAAATTTACTCGCAGATCATAGCAAGTTCAATAAGTTGATTITATACCTTTA  
 TAATGTATCAAATCTTGATACTATTGTTAGATTCTAAACTAAGTGAATCACTCTTTAAAGCTATCTAAACACA  
 35 TAAAGTCATCAAGCCTAA

ITVKQIMDEIAVSMDTARRYLQELADKDLLIRVHGAEKLRNTSLLNERSNIEKOALQTAEKQELAHFAGSLVEERETI  
 FIGPGTTLEFFARELPIDNIRVVTNSLPVFLILSERKLTDLILIGGNYRDTGAFVGTTLQNLNSLQFSKAFVSCNGIQNGA  
 LATFSEEAGEAQRIALNNSNKYLADHSKFNKFDFYTFYNVSNLDTIVSDSKLSDSILFKLSKHIVKIPZ

**ID10 483 bp**

40 ATGACTGAGTTCTGTTAGATCTCTAGAACCCATTAAACTAGCTGTTGGACCTACTACTATCACTTGAAAC  
 AGCTAGACAAAACAGATAAAAGACCAAGAGCTTAAACTGAAATTCAATCCATCTTATCGAACACAAGGGAAATT  
 45 ATGCTTATCGCCGGGTTCATTTAGAACTAAGAAATCGTTATCTGTTAAATCATAAAAGAGTTCAAGGCTTGT  
 GAAAGTACTCAATTACAAGCTAAATGCGAAAGAAACGAAATATTCTCTATAAAGGAGACGTGGTAAGAA  
 GGCAGAGAATCTCATCAAGCCCAATTGAAGGCTCTAAAACAATGAAAAGTGTACACAGATGTGACTGAATT  
 TGCCTTCCAGCAAGTACTCAAAGCTTACTTATCACCAAGTTAGATGGCTTAACAGCGAAATTATTGCTTTA  
 ATCTTCTTGTGCGCTAATTAGAATAA

50 MTEFSDLLEAIKLARWTYYHLKQLDKDQELKTEIQSIEHKGNAYRRVHIELRNRYL VNHKRVQCLMKV  
 LNLQAKMRKRYSSHKGDVGKAENLQAFEGSKTMCKYTDVTEFAIPASTQKLYLSPVLDGFNSEIIAPNLSCSPN  
 LEZ

**ID14 1266 bp**

55 CCAGGATTTGCTACCGTTGCAAGTGGTGCCTTCTCTCTAAAGGAAAATGGAGGAAAATCAATCAATCAGCA  
 CATTCAAGATCAAAGTGTCAAGGTATTGGTCAGGATGAAGATGAAAAAAATCGCTGCTGCAGCAGGGAAAT  
 GACTTTAACCTTGTAACCAATGTGGATGATATTCTACGACCAAGGATATTACTATCGTAGTGGAAATTGATGGGGC  
 60 GTATTGAGCTGCTAAACGTTTATCACTGTCGCTTGGAAAGCTGGAAAACAGGTGTTACTGCTAACAGGACCT  
 TTAGCTGTCATGGCGAGAATTGCTAGAAAATCGCTCAAGCTAACAGGTAGCACTTACTACGAGCAGCAGT  
 GCTGGTGGGAACTTCGAATTCTICGTACTTAGCAAAATTGCTGGCTTCTGATAAAATTACGCGCGTGCTTGAGTAG  
 TCAACGGAACTTCCAACCTCATGGTACCAAGATGGAAAGAAGGGCTGGTCTACGATGATGCTCTGGCGAAG  
 CACAACGTCAGGATTGCAAGAACAGCAGTACGAGCTAGATGGATTGATGCAAGCCTAACAGATGGTAA

5 TTTTGAGCCAATTGCTTGGCATGAAGATTGCCCTTGATGATGTAGGCCACAAGGGAATCCGAATATCACACC  
 AGAAGACGTAGCTGTAGCTCAAGAGCTGGTACGTAGTGAATAATGGTTGGTCTATTGAGGAAACTTCTCAGGT  
 ATTGCTGCAGAAGTAGCTCAACCTTCTACCTAAAGCGCACCCTACTGCTAGTGTGAATGGCTAATGAACGCTG  
 TCTTGAGAACATCTATCGTATTGGTAGTCTATGACTACCGACCAGGTGCGGGTCAAAAACCAACTGCAACAAAG  
 10 TGTTGAGCTGATATTGTCGATCGTGTGAGTCTATGACTACCGACCAGGTGCGGGTCAAAAACCAACTGCAACAAAG  
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 GTCAAGGTCTGAAGTTGGCTGAATCTCAATGCTCAAGATATTCCCTTAAGCAAATCCTCAAGATGGCAAAGA  
 GGGTACAAGGCGCGTGTGTTATCATCACACACAAGATTAATAAGCCAGCTGAAAATGTCAGCTGAATT  
 GAAGAAGGTTCAGAATTGACCTCTGAATACCTCAAGGTGCTAGGAGAATAA

10 PGFGTVASGVFPLLKENGKINQSAHSIDIKVAKVLVDEDEKNRLAAGNDNFVTNVDDILSDQDITIVELMGRIEP  
 AKTFITRALEAGKHVVANKDLLAVHGAELLEIAQANKVALYYEAAVAGGIPILRTLANSLASDKITRVLGVVNNTSNF  
 MVTKMVEEGWSYDDALAEAQRLGFAESDPNDVDGIDAAYKVMVLSQFAFGMKIAFFDVAHKGIRNITPEDVAVAQE  
 15 LGYVVKLVGSIETSSGIAAEVPTFLPKAHLPLASVNGVMNAVFVESIGESMYYGPAGQKPTATSVVADIVRVRRL  
 NDGTIGKDFNEYSRDLVLANPEDVKANYYFSILALDSKGQVLKLAEIFNAQDISFKQILQDGKEGDKARVVIITHINKA  
 QLENVSAELKKVSEFDLNLNTFKVLGEZ

ID16 1725 bp

20 ATGAAACACCTATTATCTTACCTCAAACCCATCAAGGAATCAATTAGCCCCCTTGTCAAGCTGTTAGAAG  
 CTGTTTTGAGCTCTTGGTCCATGGTATTGCTGGGATTGACCAATCTTACCTCAGGGAGATCAAGGTCT  
 CTCGGATCAGATTGGCCTGCTCTTATCTTGCACTGAATTGGCCTTACTGGCTGATAGCTCAATTCTACTC  
 AGCAAAGGCAGCTAGGTTCTGCTAAGGAATTGACAAACGATCTTACCTCGTATATTCTTCTTGCCCAAGGAC  
 25 AGCAGAGACCGTCTGACAACCTCTAGTTGGTCACTCGCTTACCTACCAAGATTCAGACTGGTATCA  
 ATCAATTCTGCTCTTACGCGCCATTATCGTTTGGTGCATTATGGCTTATCGAATCTCAGCT  
 GAGTTGACTTTCTGGTCTTAGTCTTGGCCATTGACCATTTGACCTTGTGATTGAGGTTATCTGATTGGTCAATCC  
 TTCTACAGTAGTCAGAAAGAAAACGGACCAACTGGTCAGGAAACGCGCCAGCAATTGCAAGGGATGCGGGT  
 30 TTTCTACAGTAGTCAGAAAGAAAACGGACCAACTGGTCAGGAAACGCGCCAGCAATTGCAAGGGATGCGGGT  
 GAAAAGACAGGTTCTGGTCTAGTTATTAAACACCTCTGACCTATCTGATTGCTAATGGAACTTCTCGTTATTAT  
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 35 TCGAGGAAGTCTTGTGAGGCTCCAGAGGATATCCATTCAAGAGTTAGAACAAAAGCAAGCTACCAAGAGATAAGG  
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 ACTCAAGGACAATTCTAGGTATCATGGGGAAACTGGTCTGGTAATCAAGCTGGTCAACTCTTACTGGAC  
 40 TTATCCAGTAGACAAGGGAAACATTGACCTTATCAAATGGACGTAGTCCCTTAATTGGAGCAGTGGCGGT  
 CAAGAAGTATCTGACCAGGAACTCTGGCAGGCCCTGGAGATTGCGCAAGCTAAGGATTGTCAGTGAAGGAA  
 GGACTCTTGATGCTCTAGTTGAGGCCAGGGGGGAAATTCTCAGGGAGACAAAAGATTGCTATCGCC  
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 45 AGCTCTTGAAGCTATTAGAGAAAATTCCAAACACGAGCTTAATTGATCTCTAACGAACCTCAACTTACA  
 GATGGCGGACCAGATTCTCTCTGGAAAAAGGTGAGTTGCTAGCTGGCAAGCAGCAACTTGATGAAATC  
 CAGCCAAGTCTATTGTGAAATCAATGCATCCCAACATGGAAAGGAGGACTAG

50 MKHLLSYFPYIKEYSILAPLFKLLEAVFELLVPMVIAGIVDQSLPQGDQGHLMQIGLLIFAVIGVLVALIAQFYSAKAA  
 VGSAKERLTDLYRHILSLPKDSRDLTTSSLVTRLTSPTYQIQTGINQFLRLFLRAPIVFGAIFMAYRISAELTFWFLVLV  
 ALTTIVVGLSLRVNPFYSSLRKKTQLVQETRQLQGMVRIRAFGQEKRELQIFQTLNQVYARLQEKTGFWSLLTPLT  
 YLIVNGTLLVIIWQGYISIQGGVLSQGALIALINYLLQILVELVKLAMLINSLNQSYISVKRIEEVFEAPEDIHSELEQKQA  
 TRDKVLQVQELTFYPDAAQPSLRYISFDMTQGQILGIIGGTGSGKSSLVQLLLGLYPVDKGNIDLQNRSPLNLEQWR  
 SWIAYVPQKVELFKGTIRSNTLGFNQEVSQELWQALEIAQAKDFVSEKEGLLDALVEAGGRNFSGGQKQLRSIARAV  
 LRQAPFLIDDDATSALDTITESKLLKAIRENFPNTSLILISQRTSTLQMADQILLKEGELLAVGKHDDLMKSSQVYCEINA  
 SQHGKEDZ

ID18 1224 bp

55 ATGAAACGTCTCTGACTCAAGAGTCGATTACAGTTGCTCTGCCAGTATTTTCTACTGGTCATCGGTGTGGT  
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 GCCTTGGGGCTTGTGATTTGGTTCTGGTCATGCTCTTAAACAGAATTCTTGGAGGTGACCCCTTCTATA  
 TATTTTAGGCTTGGGACTTATGATCTTGGCATTGTATTATCAAGCTTGTGATCAACGGTGC  
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 60 CCTCTGTCATTGTCCAATTCAAAAGAAAACAATAGGAATGGAGACGCCAGGGTCCGCTGGACTTTGTTAATT  
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 ACAGGAGTGTGCTGGTTCTAGCTATCTTATTAGCAAGGACGGAGCTTCTCACCAGATTGGAAATGCCGA  
 CCTACCAAAATTACGGATTITGGCTGGCTCAATCCCTTGAATTGCCAAACAAACGACTTACAGCAGGCTCA

AGGGCAGATTGCCATTGGGAGTGGTGGCTTATTGGTCAGGGATTTAATGCTTCGAATCTGCTTATCCCAGTTGA  
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 TTATGATGTTGCTCTTCACATCTTGAGAATATCGGTGCTGACTGGACTACTTCCTTGACGGGGATTCCCTG  
 CCTTCATTTCGCAAGGGGGATCAGCTATTACGTAATCTGATTGGTGTGGTTGCTTATCGATGAGTTACCA  
 GACTAATCTAGCTGAAGAAAAGAGCGGAAAAGTCCCATTCAAACGGAAAAGGGTTGATTAAAACAAATTAAATA  
 A

5 MKRSLLSRVDYSLLPVFFLLVIGVVAYIAVSHDYPNNILPILQQVAWIALGLVIGFVVMFLNTEFLWKVTPFLYILGL  
 10 GLMLPIVFNPSLVLASTGAKNWVSINGITLFQPSEFMKISYIILMARVIVQFTKKHKEWRRTVPLDFLIFWMILFTIPVL  
 VLLALQSDLGTALVFVIAFSGIVLLSGVSWKIIIPVFTAATGVAGFLAIFISKDGRAFLHQIGMPTYQINRILA  
 WLNPFEF AQTTTYQQAQGQIAIGSGGLFGQGFNAASNLLIPVRESMDMIFTVIaedFGFIGSVLVIALYLMILYRMLKITLKSN  
 NQFYTYI STGLIMMLFHIFENIGAVTGLLPTGIPLFISQGGSJAIISNLIGVGLLSMSYQTNLAEKSGKVPFKRKVV  
 LQIKZ

15 ID22 987 bp

ATGGTGGCTAAGAAAAAAATCTTATTGGTCTTTCTTGGAGGTGGTCAGAGAACATTCTATCAA  
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 ATCTGTTCCAAGCATGTACGCATTAAAATCCCTCAAGATTATCGCCAACAGATGGTACGAGCTTTTG  
 TGGAGAATGAGAATTATTTCCAAGACTGACTCGTGTGCTTGTAAAAGAGTATTGATGTTGAAGTTCTT  
 TACCATATTGAACTTCAAGGACTGCTTGTCTAAAAGAGAATCACATAGAAGGCCAGTGGATGCTGCA  
 ATACAATTGAGGGAAAGAACTCTTCAAGGAAGTTATCCAGATTATACCTCTAAATTACAGACAATCTAC  
 ATTTAAAAAGACCAATTCTATCAAGGAAGTTATCCAGATTATACCTCTAAATTACAGACAATCTAC  
 GATATGATTTCAGACTATTCTAGAAAATCTCAAGAGAACATCGATATCGAGATTGCTCCTCAAAGTATCTG  
 TAC TATCGGACGGATTGAGGAAAATAAGGGTCTGACCGTGTAGTGAAGTGTACGATTATTACACCAAGAGGGAA  
 AAACATATCATCTCTATTATCGGGGCTGGTGTATGGAAGAGGAACCTAAAGAGCTCAAAGAGTATGGGAT  
 TGAGGACTATGTACATTCCCTGGTATCAAAAAAACTCTATCAGTATCTATCAGCAGAAAGTCTTTGTCTA  
 TGCTAAACAAGAAGGTTCTGGAGTGTATGAGGCTTGTGACTGGGACTCCCTTATCTTACCGACGT  
 TGGAGGGCTGAGGAATTATCCAAGAAGGAGGATTGGACAATATTGAGAGCAATCAAGAGGCA  
 GCTCAGGCAGGCTTGTGACTGGGAGGCTAGCCAATTCAAGAAGTGTGATGAGGCTAGCCAATTCA  
 ACAAAACAAATCGAACAAAGTAGAAAAACTATTAGAGGAGTAG

30 MVAKKILFFMWSFSLGGGAEKILSTIVSNLDPEKYDIDILEMEHFDKGYEVPKHVRLKSLQDYRQTRWLRAFLWRM  
 RIYFPLRRLVKDDYDVEVSFTIMNPPLFSKRREVKKISWIHSIEELLKDSSKRESHRSQLDAANTIVGISKKTNSIK  
 EVYPDYTSKLQTYNGYDFQTIKEQSKEIDIEIAPQSICHTIGRIEENKSDRVEVIRLLHQEGKNYHYFIGADMEEEL  
 KKRVKEYGIEDYHFGLGYQKNPYQYLSQTKVLLSMSKQEGFPGVYVEALSLGLPFISTDVGGAELSQEGRFGQIESNQ  
 EAAQAITNYMTSASNFVDDEASQFICQQFTIKQIEQVEKLLEEZ

40 ID23 1434 bp

ATGGAAACTGCATTAATTAGTGTGATTGTGCCAGTCTATAATGTGGCAGTACCTAGAACAAATCGATAGCTTCCA  
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 CAGAGCTTATATGAGCAATTAGTCAAGAAGATGCGGATGTTGAGCTGTGTCATGAATGTCTATGCTAATG  
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 AGGTGAAAAAAATACCTGGGACGATTGCAATAAGCTAACAGAGAACATGCAACTGCCCTATCCTTCTCTAA  
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 50 ATATCTACCAAAAGTTTATGAAAGTTGTAAGGAACTATCCTGACTGAAAGAGGTCGTTTTTCAGATTGGC  
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 GGGCTATTGATAAAATATTCTTATATGATTACTGAAAATATTGAAAATCTAAAGAACATTACATTAG

55 METALISIVPVNVYAQYLEKSIASIÖKOTYQNLEILVDDGATDESGRLCD SIAEQDDRVSVLHKKNEGLSQARNDGMK  
 OAHDYLIIDSDDYHPEMIQSLEYQLVQEDADVSSCVMNVYANDESPQSANQDDYFVCDSQTFLKEYLIGEKPCTI  
 CNKLIKRQIAATPSFPKGIIYEDAYYHFDLILKAKYYVNTPKYYYYFHRGDSITTKPVAEKDLAYIDIYQKFYNEVVKN  
 YPDLKVEAFFRLAYAHFFILDKMLDDQYKQFEAYSQIHLFLKGHAFAISRNPIFRKGRRISALALFINISLYRFLLKNE  
 KSKKLHZ

60

ID24/735bp

ATGAGAATCAAAGAGAAAACCAATAATATTAAATGGAGGAATAAAAATGTAAGTAAGCATTATGGTCATCAATC  
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 AGTACGTTGATGAAAATTCTTGTTCAGAATAATCAACCGACTCAGGTATATTATAAGCAGTGATAATGGGG  
 ATTAATCGAGAACAAAATTATTITATCTAAAACAGGTAGAGAATTAAAATATTGTCAAATTATATGG  
 5 TGGTACTACAATCAAGAAAGATTAGATGTTGATCCAAGAGTTAGATTGACTCAGTCTATTAAATAAAAAGTA  
 AAGACCTATTCTGGGTACAAAACAAAAATTAGCTTCTAACTCTCGTACGGAACCTGATATAATTGATTIT  
 AGATGAACCGACTAATGGTTAGATATTGAATCATCACAAATAGTTAGCGGTTCTAAAAAAATTAGCTTACAT  
 GAAAATGTGGAATTITAATATCGACTCATAAATTAGAACAGATTGAAGAAATTGTGAGAGAGTTCTTCTGG  
 10 AGAACCGGCTTGTACATTCAAAAAGTAGGAAAAGATAGTCATAATTCTTGTGAGAGATAGCTTTCATCAGC  
 TACAGATAGAGACATTTTCACTACCAAAACAAGAATTGGATATTGTTAG

MRIKEKTNNINGGIKNVSKHYGHISIILKDINFALNKGIEVGLAGRNGVKSTLMKILVQNNQPTSGNISSDNVGYLIEEP  
 KFLSKTGLENLKYLNSLYGVVDYNQERFRCLIQEELDTQSINKVKTYSLTKQKLALLTLVTEPDILILDEPTNGLDIE  
 SSQIVLAVLKKLALHENVGILISSHKLEDIEEICERVLFLENLTFQKVGDHSNFLFIAFSSATDRDIFITKQEFWDIVZ

**ID25 1704bp**

ATGACTGAATTAGATAAACGTACCGCAGTAGCATTATGACAGCATGGTAAATCACCTAACCGTGTATGCTTC  
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 20 CATGTAACATTCACTTGCATGATTGGGAAACTGGCTAAAGAAGGTGTCAAATCTGCAGGCCGCTTGGCTGTACA  
 GTTGGACCCTTACCGTAGCGGACGGGATCGTATGGGAAACGCCGCTGTGATGCCCTCTCTAAACATCTGTGAC  
 ATCATCGCGACTCCATCGAGCGCGCTATGAGTGGTACAACCGGGATGCCCTCGCGTATGGCTGTGACA  
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 25 ATGACAGCTGAGGACGTAAACGCTTGAATGTAATGCCCTGCCCTGGCCCTGGTGGTGTGGTATGTAACTG  
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 TGATAAGAAAGAAGATATCGAAGCAGCAGGACGTGCTGTTAAGATGTTGAACTTGGTCTCAAACCCATCAGA  
 TATCTTGAECTGTGAAGCCTTGAAGATGCTACTGTAACGATGGCTCTGGTGGTCTACAAACGCCACTCTT  
 CACTTGCTGCCATTGCCCATGCCAAATGTTGACTTGTCACTGGCTCTGGTGGTCTACAGATTCAAGAACGTGTG  
 30 CTCACTTGGCCACTTGAACCATCTGGTCACTGGTCACTGGTCACTGGGACTCTCAACGATTCAAGAACGTGTG  
 TATGAAGTATTGTGCAATGGTCTTCACTGGAGATCGCATCACATGACTGGTAAGACTGTAGCTGAAAAC  
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 35 CGATGTAGTCGTTGCTGTTGACTCAGAAGAAGATGCCCTCTGGCTCTGGCTGAGATGAAATCGTGTG  
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 40 CCACCACTTACAGCCGGTGTCTCGTAAATATGCCACATCGTATCATCTGCTTACGCCGAGCCGTGACAG  
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MTELDRHRSSIYDSMVKSPNRAMLRATGMDKDFETSIVGVISTWAENTPCNIHLHDFGKLAKEGVKSAGAWPVQFG  
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 45 KIDILVSVFEGIKWNHGDMDTAEDVKRLECNACPGPGCCGMYANTMATAIEVLGMSLPGSSHPAESADKKEDIEA  
 AGRAVVKMLELGLKPSDILTREAFEDAITVTMALGGSTNATLHLLAIAHAANVDSLSEDFTIQERVPHLADLKPSQY  
 VFQDLYEVGGVPAVMKYLLANGFLHGDRITCTGTVVAENLADFADLTPGQKVIMPLENPKRADGPLIILNGNLPDGA  
 VAKVSGVKVRRHVGPAKFSEEDAIQAVLTDEIVDGVVVVRVGPKGPGPMPEMLSLSMIV/GKGQGDKVALLTD  
 GRFSGGTYGLVVGHIAPEAQDGGPIAYLRTGDIVTDQDTKEISMAVSEEELKRKAETLPPLYSRGVLGKYAHIVSSA  
 SRGAVTDFWNMDKSGKKZ

50

**ID26 274bp**

ATGTTATAATAAAATAAGAATTAAAGGAGAAATACAATATGTCATTGGAGGAGCATGGCCATATGC  
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 55 GGAGAGGAAGTTTATGTTCTGGAGTATTGTAATGGAAACCCCTATTCTATCAGAGCTAAAAAGAAAAT  
 AAGTCTGTGAAAGAAATTGCTGATTITATCATAAGGAATTAAATCCA

60

CYNKNKEFKEKYNMSIFIGGAWPYANGSLHIGHAAALLPGDILARYYRQKGEELYVSGDCNGTPISIRAKKENKSVK  
 ELADFYHKEFNP

**ID28 1065bp**

ATGACAACATTATTCAAAAATAAGAAGTAACAGAACCTGCTGCAGTCAGGTATGAAGGCCCTGTCCTG  
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1 ACATTAGAAGCTGTGGATGCACCGCGCTTGGCGCTCATATGGACGAAGTTGGTTATGGTCAGCGAA  
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 AACTCTTGACTCGTATGGTCACTGAAATTCTGTGATTTGATTCAGGTTCTGTCCTCCGCACTTGACTCGTGGAAAGGG  
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 5 GGCATCCCGTCTGGTATACCGATACCTGTCAGGATAGTCTGCAATTGACAGGCAATGAAAAAAATATCATCTCAA  
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 10 AACCTGATTCTGTTCTATGATCCAGGTCACTTGCTTCTCCAGGGATGAAGGAATTCTCTTGTACAACGGTCAA  
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 TCCTAGAAGCGCAAGCTTCTACAAGCCTGGTGAAGAAATTGGATGTTCAACGGTTGATTGATTAACATTA  
 TTAA  
 15 MTTLFSKIKEVTELAAVSGHEAPVRAYLREKLPHDEVVTDTGLGGIFGIKHSEAVDAPRVLVASHMDEVGFMVSEIKP  
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 VPDSSAITANEKNIISKAWDNRYGVLMVSELAEALSGQKLGNEYLGSNVQEEVGLRGAHTSTTKFDPEVFLAVDCSP  
 AGDVYGGQKGKIGDGTIRFYDPGHLLPGMKDFLLTTAEEAGIKYQYYCGKGGTDAGAAHLKNGGPSTTIVCARYI  
 HSHQTLYAMDDFLEAQFLQALVKKLDRTVDLIKHZ  
 20 **ID31 1182bp**  
 ATGGAATTCTATGAAATCACTGAAAGGACTACTCTTATCATAGCTAGTTTATCTGACTCTTGACTTGGAT  
 25 GAACACTTCTCCCATTCACTGATTCCAGGACTAGCTTAAACAAGCCTATCTGACTTTATCTCTAGCCACTCGTC  
 TCCCACTACTAGAAAGCTGGTTACAGTTGGAGAAGGTCTACCCGTCACAAATTACAGCCTTCTCTCAAT  
 CATCCTACTAACTCTTCAACTTATGTTGGGGCTCTCGCTTAGCTGTCAGTTGGCAATCTG  
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 TCACCCGCTGGTTACCTAGCCTATTTAGGACTCTTACATGATAATGGGCAATCGTCTCTTACAT  
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 30 AAAAGATTCTTCTCCCCTATCTAGGAAAATTACCCATCTCAAACGTTAAATCACGATACTAGAGAAATTCAAAT  
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 GCTCCGCATCCCTTTCTATCTCAGGAGGTCTGGTCAAACCTTTACTTACTGTTAAACTCAGGCGACCATAC  
 CAAGAATATCTATGATAATCTCAAGCCGGCAGCAAAGTAAACCTAGACAGAGCTTACGGACACATGATCATAGA  
 35 AGAAGGACGAAAATCAGGTTGGATTGCTGGAGGATTGGGATCACCCCTCATCTTACATCGTGAACAT  
 CCTATTAGATAAAACAGGTTCACTTACTATAGCTTCCGGAGATGAAAATGCACTACCTAGATTACTCC  
 GTAATGCTCAGAAAATCTAATTGAACTCCATCTAATGACAGTACGAAAGACGGCTATCTTAATTGAA  
 ACAAAAAGAAGTGGCCGAACATGCAACCGTCTATATGTGGCTCTATTCTATGATGAAGGACTTGCCAAACA  
 GATTAAGAAAACAAATCCAAAACAGAGCATATTAC  
 40 MEFSMKSVKGLLFIAASFILTLLTWMNTSPQFMIPGLALTSLTFILATRLPLLESWFHSLEKVTVHKFTAFLSIILIFH  
 NFSMGGLWGSRLAQQFGNLAIYIFASILYAYLGKYIQYEAWRWIHLRYVLYAYILGLFHTYMIMGNRLLTFNLLSFLVGS  
 YAEGLLAGEYIIFLYQKISPYLGKITHLKRNLNDTREIQHLSRPFNYQSGQFAFLKIFQEGFESAPHFISGGHGQTLV  
 PTVKTSGDHTKNIYDNLQAGSKVTLDRAYGHMIEEGRENQVWIAGGIGTPFISYIREHPILDQVHFYYSFRGDENAV  
 YLDDLRLNYAQKNPNFELHLLIDSTKDGYLNFEQKEVPEHATVYMCGPISMKALAKQIKKQNPKTEHIY  
 45 **ID32 900bp**  
 ATGACTTTAAATCAGGCTTGTAGCCATTAGGACGTCCAATGTTGGAGTCAACCTTTAAATCACGTTAT  
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 50 TAAGGAGCAAATTGCTTATGACACACCAGGGATTACAAGCCTAAAACAGCTCTCGGAGATTCTGGTTGA  
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 GACGATATGATATGAGCGCTCTAAGGCTGCAAGGTTCTGTGATTGGTTGTAATAAAATCGATAAGGTCC  
 ATCCAGACCAAGCTCTGTCAGATGATGACTTCCGTAATCAAATGCACTTTAAGGAAATTCTCAGC  
 CCTCTGAGGGAAATAACGGTCTGCTGCTAGTGGATAATTGAGTGGAAATCTGGATGAAGGTTCCAATATTCTCCG  
 55 TCTGATCAAATCACAGACCCATCCAGAAACGTTCTGGTTCAAGAAATGGTCTCGGAGAAAAGTCTGGCACCTAACFC  
 CTGAAGAGATGCCCATCTGTGAGGAGTGTGACTCTATGAAACGAGACGAGAGACAGACAAGGTTACAC  
 TCGTGCACCATCAAGTCAGGAGCCGATAGCAAAAGGGATTATCATGGTAAAGGTGGCGCTATGCTTAAGA  
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 TCAAGAAAACAGGCGATAAAAGCTAGATTGGCTGACTTGGCTATAATGAAAGAGAATCTAA  
 60 MTEKSGFVAIILGRPNVGKSTFLNHVMQKIAIMSDEKAQITRKIMGIYITDKEQIVFIDTPGIHKPKTALGDFMVESAYS  
 TLREVDTVLFMVPADEARKGDDMIERLKAAYPVILVVNKIDKVHPDQLLSQIDDFRNQMDFEKEIVPISALQGNVVS  
 REVDSLSENEDEGQYFPSDQYDHPERFLVSEMVRKVLHLTREIPIHSVAVVVDMSKRDEETDKVHIRATIMVERDSQ  
 KGIIIGGGAMLKKIGSMARRDIELMLGDKVFLETWVKVKKNRDKLADFGYNEREYZ

ID33 855bp

5 CTGCTTCTTGTACAGAAGGAGGACTTATGCCTGAATTACCTGAGGTTGAAACCGTTGTCGTGGCTAGAAA  
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 10 TTCAAAGGGAATTGCCTAGTCAGATTATCGAGTCATGGGACGTGTAAGGAAATTTGCTTCTGACAGA  
 CAAGGTCTTGATTCCCATTGCGGATGGAGGGCAAGTATTACTATCCAGACCAAGGACCTGAACGCAAGCAT  
 GCCCATTTCTTCATTGAAAGATGGTGGCACGCTGTTATGAGGATCTCGCAAGTTGGAACCATGGAAC  
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 GCTGCTTGGCAATATCTATGTGGATGAGGTTCTGGCAGCTCAGGTTCATCCAGCTAGACCTCCCAGACTT  
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 TGGTCAAGAATGTGTACGCTGTGGTACCATCATTGAGAAAATTCAACTAGGCGGACGTGGAACCCACTTTGTCCA  
 15 AACTGTCAAAGGAGGGACTGA  
 MLLVFTEGGLMPPEVETVRGLEKLIIGKKISSIEIRYPKMIKTDLEEFQRELSQIESMGRRGKYLLFYLDKVLSHL  
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 20 KSKKPIKSHLLDQTLVAGLNIYVDEVLWRAQVHPARPSQLTAAEATAIHDTIAVLGQAVEKGSTIRTYTNAGED  
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ID34 633bp

25 TTGTCCAAACTGTCAAGGAGGAGCTGATGGGAAAAATCATCGGAATCACTGGGAAATTGCTCTGGTAAGTC  
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 GCCCTCTCTAGCTAGTCTCATCTTCAAATCTGTGAACGAGAATGGTCTAAGCAAATTCAAGGGGAGATTAT  
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 30 TTGAGCAGGACTACACCGATTGGCTGAGACTGGTGGCTATGTGGACGAGTGGCTCTGGCAGCCAGTGGCCTTAGAAAAAAAGA  
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 GGGAGGTAGGCAAGATGACAGAGATTAA  
 35 MSKLSKEGLMGKIIGITGGIASGKSTVTNFLRQQGFQVVADAVVHQLOKPGGRLFEALVQHFGQEIILENGELNRPLLA  
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ID35 1269bp

40 TTGATAATAATGGCAATCAGAACAGCTTCTTAATCAAGTCATATCCTTGTAGGGAGGTAGGCAAGATGACA  
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 TACCTTATGCCCCTCTCGTGGAAAATCTAGGTGTAGGGAGTCAGCAAGTCGCTTCTGAGGCTTAGCAAT  
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 45 CTTCTCTGTTACTAACGGTGATTTGAGGTTCTAATGCAACGGCACTGATAGCCAGTCAGGTTCCAA  
 AGGAGAAAATCAGGCTCTGCCCTAGGTACTTTGCTCTACAGGCGTAGTGCAGGACTCTAAGTGGTCCCTTATTGG  
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 50 TCGGTTAAATATCCCTATCTTGTCTCAATCTCTTAAACAGTTGTCTCATCCAATTTCAGCTCAATCGATTGG  
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 55 CTGCAGTAGCAGGTCAATTGGCAACCATGCTGTCTTATGCGACAAGCCTTGTGTTGCCCTTAGTTGTCTCTT  
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60 MIIMALRTSFLKCISFLKEVGMTEINWKENLRIAWFGNLTGASISLVVPFMPIFVENLGVGSQVAFYAGLAISVSAIS  
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 FVIQFSAQSIQGIPILALYVRDLGQTENLLFVSGLIVSSMGFSSMSAGVMGKLGDVKVNHRLLVVAQFYSVIIYLLCANAS  
 SPLQLGLYRFLFGLGTGALIPGVNA!!SKMTPKAGISRVAFNQVTFYLGGVVGPMAVGQFGYHAVFYATSLCV  
 AFSCLFNLIQFRLLKVKEIZ

ID36 131bp

ATGGCCCTACCAACTATTGCCATTGAGACGCCAATGTTGGAAATCAACCCATTAAATCGGATCGCTGGTG  
 AGCGAACATCTCATTGAGAAGATGTCGAAGGAGTGACACGTGACCGTATTTATGCAACGGGTGAGTGGCTCAATC  
 5 GTTCTTTAGCATGATTGATAACAGGAGGAATTGATGATGTCATGCCCTTCATGGAACAAATCAAGCACCGAC  
 AGAAAATTGCCATGGAAGAACAGATGTTATCGTTTGTCTGGTAAGGAAGGAATTACTGATGCAAGCAGAC  
 ATACGTAGCTGTAAGCTTATAAGACCCACAACCGATTACCTCGCAGTCACAAAGGTGGACAACCCCTGAGAT  
 10 GAGAAATGATATATATGATTCTATGCTCTCGGTTGGGTGAACCATTGCTATCTCATCTGTCCATGGAATCGGT  
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 TTAGCTGATGGCTCTAACGTTGGAAAATCAAGCTGATCAATGCTATCTGGAGAAGACCGTGTATTG  
 15 CTAGCTCTGTTGCTGGAACAACCGTGTGATGCCATTGATACCCACTTACAGATAACAGATGGTCAAGAGATTACCAT  
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 20 ATCAGCTTAACCAAGCAACGTCCTCCAAACTCTCTGAGATGATTAAGCAAATCAGCGAAAGTCAAATACACG  
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 ACGTCTCAAGATTTCTATGCGACCCAAGTGGCAACCAAACCCACCAACCTTGTCACTTTGTCATGAAGAAGAA  
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 25 TCTATCGCAAGAAAACGCAAATAA  
 MALPTIAIVGRPNVGKSTLFNRIAGERISTVEDVEGVTRDRIYATGEWLNRSFSMIDTGGIDDVDAPFMEQIKHQAEIAME  
 EADIVFVVSKEGITDADEYVARKLYKTHKPVILA  
 30 VNKVDPNPEMRNDIYDFYALGLGEPLPISSVHIGTGVDVLDIAVE  
 NLPEYEEENPDVIFKSLIGRPNVGKSSLINAILGEDRVIASPVAGTTRDAIDHTFTDTDGQEFTMIDTAGMRKSGKVE  
 35 NTEKYSVMRAMRAIDRSVVLMVINAEGIREYDKRIAGFAHEAGKGMIVVNWKWDTLEKDHNMTKNWEEDIREQFQ  
 YLPYAPIIFVSALTQRLHLPEMIKQISESQNTRIPSAVLNDVIMDAIAINPTPTDKGKRLKIFYATQVATKPPTFVIFVNE  
 EELMHFSYLRLENQIRKA  
 FEGTPHI  
 LARKRKZ

**ID37 714bp**

30 ATGACAGAAACCATTAAATTGATGAAGGCTCATCTCAGTGCAGGTTAAAGAGCAAGAAATTCCCCAAGTA  
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 TCTCTTGTGGAGATTGAGCAGAAAAGGGAGCCGACTTCATACCGACACCTTCAACCCCAAGGTGT  
 35 GGAAGGTCTCTTGTGATTAGTCGGTCATGCGAGCTCTGCTGACAAAAGCCTTGTGGCAGCTGAAAGCTTGGGC  
 TATGGTGGTGTGATTATCGGTTGGTGTGACATAAGCTGAAGAAGTGGCAGAGCTTTAACCTACCTGACTACA  
 CCTATTCTGCTTGGTGTGACACTGGGTGTGCAATCAACATCATGATATGAAACCGAGACTGCCACTAGAGAA  
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 GCTGGGGCGCGTGCACCAAGCTGGAGTCAGCGCCTAGCAGAACAGTTGGTCAAGCTGAACCAAGCTCAACT  
 40 AGAAAAAAATCTGAACAGAAGAAATTATTGTAG  
 MTEETIKLMAHTSVRRFKEQEIPQVDLNEILTAQMSSWKNFQSYSVIVRSQEKKDATYYELVPQEAIRQSAVFLLFV  
 GDLNRRAEKGARLHDTDFQPGVEGLLISSVDAALAGQNLAESLGYGVIIGLVRYKSEEVAFNLPDYTVSVFG  
 45 MALGPVNQHDMKPRPLENVFEEEYEQSTEIQAYDRVQADYAGARATTSWSQRQAEQFGQAEPSTRKNLEQK  
 KLLZ

**ID38 729bp**

50 ATGACAGAAATTAGACTAGAGCAGCTCAGTTATGCCATTGGTCAGGAAGGATTAGAGGATATCAACCTACAG  
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 GCACAAAGGATCTGCTTGGAGCACAAGACGGTCTGGAAATATCATTCTGCCCTCTGATTCAAAAGGTGGAT  
 AAGGCAGAACGCTATTCCCGAGCGGATAAAATTCTTGCACCTCCAGCTGACAGCTGAAAGACAAAGTACCT  
 CATGAACTTAGCGGTGGATGCCAGCGTGTAGCCTACTCEGGACCTACCTTGGCACAAGCTCTCTCT  
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 55 GAAAAATCGGCTGGCAGATTGTTGAGAAATTAAACTAGATTGGTGTGAAGATGAGGACAAGGAAGTCCAAA  
 GATTGCCAACAGTCAAATTGGCGGAATTGGCTTAGATAAGTAG  
 MTEIRLEHSVAYGQERILEDINLQVTSGEVVSILGPSVGKTLFNLIAGILEVQSGRIVLDGEENPKGRVSYMLQKDLL  
 LEHKTVLGNHPLLIQKVDAEAISRADKILATEQOLTAVRDKYPHELSGGMRQRVALRTYLEFGHKLFLDEAFSALDE  
 60 MTKMELHAWYLEIHKQLQETTIIHSIEEAALNLSDRYILKNRPGQIVSEIKLDWSEDEDKEVQKLA  
 YKQQLAELGLDK

**ID39 2433bp**

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 5 GTATGAGATGGACCGTTAGAAGAGGGCTTGGACTGACTGAAACGGACTATAGCCAGGATGAAACCTTAC  
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 10 CTGGACTCGTAAGATCTAAGGCTTACGCCAACGCCATCTACAGTAGCTGACAGCAGCAGGAAATTCTATGCCAA  
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 15 GGCAGAAGTAAAGCAGTCATCGGTCGGAGGAAAATCTACGTATGATTCAAATCTTGAGCCAGAGACTAAG  
 AACAAACCTGTCTGGTGGGGATGCTGGTCTGGAAAACAGCTCTGGCCTGGCTTGCCCAGCGTATTGCTA  
 GTGGTGACGTGCTCGGAAATGGCTAAGATGCCGTGTTAGAACCTGATTGATGAATGTCGTTGCAAGGACAC  
 GCTTCGTGGTACTTGAAGAACGATGAATAATATCATCAAGGATATTGAAGAAGATGGCAAGTCATCTCTT  
 20 TATCGATGAACCTCACACCATCATGGTTCTGGTAGCGGGATTGATTCGACTCTGGATGCGGCAATACTTGAAA  
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 25 GCAAAATAAGGAAAGCATGTAAGCAGACGATTGAGCTAGATTGAGTCAGCTGACAAGGCCATGATGGATGGAA  
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 TATTTGACCACCTTGAGTCGTTGTCAAGGAAATCCAGTTCAAAACTGACTCAAACGGATGCTAAGAAGTATT  
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 30 GCAACCAGTCAGGGATTGCACTGATAAGCTGGGATTGGCTTGTGAGGATTTGACAGCTTCAAGGCTTACAGGTGCTGGAA  
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 40 AGCCTTGTGCAAGTTGACTGAAAGAACGGATTGACTGAAACATTACAAGCTCAGCTGAAACATTGTTAGCAA  
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 TGATATTGCTAA  
 45 MNYSKALNECIESAYMVAHGFGARYLESWHILLIAMSNSYSVAGATLNDYPYEMDRLEEVALELTETDYSQDETFT  
 LPFSRRLQVLFDEAEYVASVVAHVKLGTEHVLYAILHDSNALATRILERAGFSYEDKKDQVKIAIRRNLEERAGWTR  
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 GSGSGIDSTJDAANILKPALARGTLRTVGATTQEEYQKHIEKDAALSRRFAKVYTIEPSVADSMILQGLKATYEHHRV  
 QITDEAVETAVKMAHRYLTSRHPDSAIDLDEAAATVQNKAHKVAKDSDLSPADKALMDGKWKQAAQLIAEVEV  
 PYYKDLVTESDILTLSRLSGIPVQKLQTDAKKYLNLAEELHKRVIGQDQAWSISRAIRRNQSGIRSHKRPISGMFLGP  
 TVGVKTELAKALAELVLFDDESALIRFDMSYEYMEKAFAASRLNANGPPGVYVEEGGELTEKVRNPKPSVLLFDEVEKAHP  
 DIFNVLQVLLDDGVLTDSKGRKVDFSNLIMTSNLGATALRDKTVFGAKDIRFDQENMEKRMFEELKKAYRPEFIN  
 RIDEKVVVFHSLSSDHMQEVVKIMVKPLVASLTEKGIDLKQASALKLLANQGYDPEM GARPLRRTLQTEVEDKLAELL  
 KGDLVAGSTLKIVKAGQLKFIAZ

**ID40 1008bp**

50 ATGAAGAAAACATGGAAAGTGTAAAAACGCTTGTAAACAGCTCTGTAGCTGTTGCTGGCCTGTGGTCAAG  
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 ACCACACAGGGCTTATGTTGCCAGGAAAAGGTTATTCAAAAGAAGCTGGAGGGATGTTGAATTGAAATTGC  
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 55 TCGTAAATCTGATAATGTAAGCAGTCAAAAGACTTGGTGGTAAGAAAATATGGACATGGAAATGACCCAACTGA  
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 60 AACGTGACTTGTGATCGAATCTAAAAATACTGTCAAAAGAATACGCAAGCAGACAAGGAAAATGGTATCCTTAAAGAAGACTTGACAGACA  
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 AAGGCTGACCAACGAATTGTGAAATAA

MKKTWKVFLTVTALVAVVLVACGQGTASKDNKEAEKKVDFILDWTPNTNHTGLYVAKEKGYFKEAGVDVDLKLP  
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 KTLVESQGGDFEKVEKVPNNDNSNITPIANGVFTAIIYVGWDGILAKSQGVANFMYLKDYVKEFDYSPVIANND  
 YLKDNEEARKVIQAIIKGQYAMEHPEEAADILIKNAPELKEKRDVFIESQKYLKEYASDKEKWGQFDAARWNASY  
 5 KWDKENGILKEDETDKGFTNEFKZ

ID41 762bp

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 10 AGTAGCAGGTTCTTAAACTTCTCCCAAGTTATCCTGCCACACCTCTGAAATTCTCCAGCCCTTGTGCG  
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 15 CAGACCATTCCGACCATTGCCATAGCTCCTATCCTGGCTTGTGCTAGGTTATGGGATTGGCCAAGATTGCTT  
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 20 AGGCTTGGTGTATATGATTCACTGTTAGTATGATACCATGTTGCCATTATTCTGGTGT  
 CGATTATCAGCTTGGTATGAAGCTGGTCGATATCAGTGAAGAAATATGTAATTAAATGAAACGTTGCG  
 YVIKWRSZ

ID42 372bp

TTGATTTTAATCCTATTGCTGTATGATAAGGGAAAAGAAAGGGGACAGAGATATGGCTTTACCAATACCCACA  
 25 TGCGATCTGCTAGTTTGGTATTGTTACAGCTTGCCTGATGACATCATTGACTTTGGTATATCATGACCAT  
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 35 HLPTAIDFDFNHPFDPRPPRVLVDMDGREILLPEENDLFZ

ID43 1569bp

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 CAATTAGAACAGAACAGTTGAATTCACTGTTAACATCTACTAATGAAAATAAGAGTTAAGGAAGATGTTATA  
 45 AGTGACAGAAATTCAAAAAGAATTGAAGATACTGCTTAAAGTGTAAAAGATTATGGTCAGTAGGTGATGGG  
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5 ID44 324bp

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10 CAAAAGTTAGAAAAAAATCAATGGATAAGAGCGACATGCCCGTCGCAGATGGTCCAGATCGGAAGTATT  
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15 ID45 816bp

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30 MKKMKYEEETSALLHEFSEENQYFEELWESFNLAGFLYDEDYLREQIYLMMLDFSEAERDGMSAEDYLGKNPKIM  
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35 ID46 348bp

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45 ID47 1260bp

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5 MQNLKFAFSSIMAHKMRSLLTMIIGVSSVVIMALGDSLSRQVNKDMTKSQKNISVFFSPKKSKDGSFTQKQSAFTVS  
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ID48 705bp

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ID49 1200bp

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35 GTGATGCGCGTGCAGATGCTGCGGCAATTAGCAAGGCTCAAAGTCATTGGATGCAACAATGTTCTAGTA  
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50 VKADEATNZ

ID50-759bp

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5 MSRKPFIAGNWKMKNPPEAKFVEAVASKLPSLDLVEAGIAAPALDLTTVLAVALAKGSNLKVAACQNCYFENAGAFTG  
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ID51 1473bp

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30 MKTKIGLASICLLGLATSHVAANETEVAKTSQDTTASSSEQNQNSNKQTSAEVQTNAAHWDGDYYVKDDGSKAQ  
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ID52 774bp

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ID59 1071bp

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 15 QMLEKNENLRYVVPTEASNLWFDNMVPKTVKNQNSAYAFINFMLKPENALQNAEYVGYSTPNLPAKELLPEETKED  
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ID61 1851bp

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45 MNKKLTDYVIDLVEILNKQQKQVFWGIFDIFSMVVSIIVSYIILYGLINPAPVDYIIFTSLAFLYQLMIGFWGLNASIRY  
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ID101 1338bp

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 10 ACGGTGCACAATAGACTTATATCGCTCCAGACCATCTATTGAAAGTAGTGAAGTTGCTGCTTGGTTGAGACCA  
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**ID102 1512bp**

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40 MTIYNINLGIGWASSGvEY AQAYRAGvFRKLNLSSKFIFTDMILADNIQhLTANIGFDDNQVIWLynHFTDJKIAP1SVT  
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 45 SQGRKPFSLITASRLAKEKHIDWLVKAVIEAHKELPELTFDIYGS GGEDSLLREIIANHQAEDYIQLKGHAELS QIYSQYE  
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50

**ID103 2292bp**

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**ID104 879bp**  
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**ID106 327bp**  
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ID108 954bp

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ID110 1902bp

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ID111 1179bp

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ID113 2466bp  
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ID114 1974bp

10 ATGAAAAAAATTITATGTAAGTCCAATTITCCTATTCTAGTAGGATTGATTGCGTTGGAGTCCTATCCACTTTCAT  
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ID115 663bp

50 ATGAAGTCCTGTTATGCGGAGACTATGAAGACTGTTAACCTTCTAGTCCTACTCTGAGGAATGATG  
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60 MECLICGQTMKJVLTFSSLLRNDDSCI.CEDCDSTFERIGEENCPNCMKTELSTKCQDCOLWCKEVSHRAIFTY  
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ID116 1299bp

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 20 GGCTGGTCTATGA

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ID117 870bp

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 45 DYPETQFYEPHRVADTLLENMILEVYGDRSVVLVRELTKIYEYQRGTISELLESIAETLKGECLLIVEGASQGVEEKDE  
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ID118 345bp

50 ATGATAAAAGAAAGGAAAGGGCTTTATGGACAACAAAGAATTATITGACGCGCTGGATGATTCTTCCAACAA  
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 55 MIKKGKGFMDKKEFLDALDDFSQQLEVTLADVEAIKKNLKSLMEENTARLENSKIRERLGVEADAPVKAHVRES  
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ID119 639bp

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ID120 408bp

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20

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ID121 285bp  
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30

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35

ID124 1311bp

40

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55

MKSRYKETSMOKIVVQGGDNRLVGSVTIEGAKNAVLPLLAATILASEGKTVLQNVPLSDVFIMNQVVGGLNAKVDFD  
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ID125 1101bp

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ID126 1281bp

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ID127 894bp

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5 VGRIIIRAGVKMELGKVFRFRSGNSLKEAAGESCSTSSQLSRFELGESDLAVSRFFEILDNIHVTIENFMDKARNFHN  
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**TABLE 3****ID1 1068bp**

5 ATGTCTAACATCAAACATGTCCTGGAGGACATCATGGGAGAGCGTTGGTCGTAECTCAAGTACATTATTCAAGGA  
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CACGTAACAGGTTCTGGACGGAGATCCTCGGGCTATGCGTTACTGAGGGACGTTGCTGAAATTG  
10 CAGGCTACCTCTTCAGGATATCGAGAAAAGACAGTCTTGCATGAACTTACGATACGGAGAAAAGAAC  
CAACGGCTTGCAGCAGCCTTCCAACCTCTGGTCAATGGTCAGTGGGATTTCGGCTGGTATGCCACAGA  
CATTCTCCCCATAATTAGCTGAGGTCAAGATGCTGCAGTTACATGATTGACCACCCAACGCAAAAGATTGAT  
AAACTCATGGAATTCTTGCTGGACCAGACTTCCACAGGGCTATTATCAGGGTGTGATGAAATCAAGAAA  
GCTTATGAGACTGGGAAAGGGCGCGTGGTGTGTTCAAGACTGAAAATTGAAAAGCTAAAAGGTGGAAGGAA  
CAAATGTTATTATTGAGATTCTTATGAAATCAATAAGGCCAATCTAGTCAGAAAATCGATGATGTTGTTA  
ATAACAAGGTAGCTGGGATTGCTGAGGTCTGATGAGTCAGCTGATGGTCTCGTATCGTATCGAACTTAA  
GAAAGACGCTAAACTGAGCTTCTCAACTACTTTAAGTACACCGACCTACAAATCAACTACAACCTTAAT  
ATGGTGGCATTGACAATTTCACACCTCGTCAGGTTGGATTGTTCCAATCTGTCTAGCTATATCGTCACCGTCG  
20 AGAAGTGA  
25 MSNIQNMSLEDIMGERFGRYSKYIIQDRALPDIRDGLKPVQRRLYSMNKDSNTFDKSYRKSASKVGNIMGNFPHGDSS  
IYDAMVRMSQNWKNREILVEMHGNNSMDGPAAAMRYTEARLSEIAGYLLQDIEKKTVFAWNFDDEKEPTVLP  
AFPNLLVNGSTGISAGYATDIPPHNLAEVIDAAYMIDHPTAKIDKLMFLPGPDFPTGAIQGRDEIKKAYETGKGRVV  
VRSKTEIEKLKGKQEIQVIIIEPYEINKANLVKKIDDVRVNNKVAGIAEVRDSDRGLRIAIELKDDANTELVNYLFKY  
TDLQINYNFNMVAIDNFTPRQVGLFQSCLASLTVEKZ

**ID12 684bp**

30 ATGCCGACATTAGAAAATAGCACAACAAACTGGAGTTCAATTAGAAGGCAGAAGAATTACATGCCCTGTG  
ACAATATACAGTTGAGCGGAGATAAACTAAAAGTAATTCCTTACTCTGTTAACCTGGGAAAGGAAAACA  
ACTACTTCCATAAAATAGCATGGCTTGCCTGAGGTATAAAACTCTTGTGATGGCGATACTCGAA  
ATTCACTGTTAGGATTAAATCTGTGAAAAAAATTACAGGGCTAACAGAAATTCTGACAGCTGA  
TTATCTACGGTTATGTGATACAAATATTGAAAATTATTGTTAGTTCAATCGGATCTGTATCACCAAACCC  
35 CAGCCTGTTACAAAGTAAAATTAAATGATATGATTGAAACATTGCTAACATTTGATTATATCATTATTGAT  
ACACCGCCTATTGAAATTGTTATTGATGCGGCAATTACTCAAAGTGTGATGCGTCCATCTGGTAACAGCAA  
CAGGTGAGGCGAATAAACGTTGATATCCAAAAGCGAACACAATTAAAACACAGGGAAACTGTTCTAGGA  
40 GTTGTGTTAAATAATTGATATCTGGTTAACAGTATGGAGTTACGGTTCTATGGAAATTATGGTAAAAAA  
AA

MPTLEIAQK~~KLEFIKKAEFFYYNALCTNIQLSGDKLKVISYTSVPGEVKTTISINIAWSFARAGYKTLIDGDTRNSVML~~  
GVFKSREKITGLTEFLSGTADLSHGLCDTN~~IENLFVVQSGSVSPNPTALLOQSKNFNDMIETLRKYFDYIIDTPPIGIVIDAA~~  
~~III~~TQKCDASILVTATGEANKRD~~I~~QKAKQQLKQTGKFLGVVLNKLDISVN~~KYGVYGSYGN~~Y~~GKZ~~

**ID13 1182bp**

45 ATGGAGGCAAATATGAAACATCTA~~AAA~~ACATTTACAAAATGGTTCAATTATTAGTCGTTATCGTCATTAGCT  
TTTTAGTGGAGCCTGGTAGTTTCAATAACTCAACTA~~AA~~ACTCAAAAGTAGTGTAAACAACACTCTAACACAA  
TAGTACTATTACACAAACTGCCATAAGAACGAAAATTCAACACAGGCTGTTAACAAAGTAAAGATGCTG  
TGTTC~~T~~TTATTACTTATTGCCAACAGACAAAATGCTGTTAGGTTCTGAATATGATGACTACTGACACAGATTCTAG  
CGAATCTCTAGTGAAGGACTGGAGTTATTAAAAGAATGATAAAAGCTTACATCGTCACCAACAATCAC  
50 GTTATTAAATGCCAACAAAGTAGATATTGCTGCTGAAATCTCTTCAGAAAAGTGAACACAGTAGCTGAGTTGGTGAATTCTA  
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CAAACGTGAACTGCTATTACCCAGTAACCTGGGGCCACTGATCAATATTCAACGGCAGGTTATCGGAATT  
CCTCAAGTAAAATTGCTACAAATGGAGGAACATCTGTTAGAAGGTC~~TTGG~~GAATTCGTC~~AA~~ATGCTGTTAT  
60 CAAATTTATTGAA~~ACAGT~~AGAAAAACGGAAAGTGA~~CCG~~GTCCAGCTTGGGAATTCGTC~~AA~~ATGCTGTTAT  
TAATGTTAGTACAAGCGACATGAGAAGACTCAATATTGCAAGTAATTGTTACATCTGGTGAATTGTTGTTGCTGCTG  
CAAAGTAATATGCCCTGCCATGGTCACTTGA~~AAA~~ATGCTGTAATTACAAAAGTAGATGACAAGAGATTGCT  
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GGAAAAGAAGAAA~~ACTAC~~CTCTATCAAAC~~AA~~AGTTCAGGTGATTAGAATCTAA  
MEANMKHLKTFYKKWFQQLVVIVISFFSGALGSFSITQLTQKSSVNNNNNNSTITQAYKNENSTQAVNKVD~~A~~VVS  
ITYSANRQNSVFGNDDTDSDQRISSEGVYKKNDKEAYIVTNH~~H~~INGASKVDIRLSDGT~~K~~V~~P~~GEVGADTFSDIAVV

KISSEKVTVAEFGDSSKLTVGETAIAIGSPLGSEYANTVTQGIVSSLNRNVSLKSEDGQAISTKAIQTDATAPGNSSGGPLI  
NIQQVIGITSSKIATNGGTVEGLGFAIPANDAINIEQLEKNGKVTRPALGIQMVNLSNVSTSDIRRLNIPSNTSGVIVR  
SVQSNMPANGHLEKYDVITVKVDDKEIASSTDLQSALYNHSIGDTIKITYRNKGKEETTSIKLNKSSGDLESZ

5 ID15 939bp

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AAACAGTGTGGCTACCTAAATGGTCAGTCAGAACGACCAATTACAGCTCAAGGAAACAGACCATGAG  
AAACGGTCCAAGTGATTACGATGAGAAGGAAGTGACTCAGAGAGATTTACTTTATTATTCGAGTATCGA  
TCCTCTATCTATCAATACAACAAGGAATGACCCTGGCTGCAATATCGAAGTGGGATTTATTATCAGGATGAAGCA  
GATTGCCAGCTATCTACAGTGGTCAGGACAGGAACCGATCTGGTCAGAAGATTGAGTAGAAGTGGAG  
CAATTACGCCACTACATTCTGGCTGAAGACTACACCAAGACTATCTCAGGAAGAACCTCAGGTTACTGTCATA  
TCGATGTGACCGATGCTGATAAGCATTGATTGATGCAGCAAACATGAAAAGCCTAGTCAAGAGGTGTGAAGG  
CCAGTCTATCTGAAGAGTCTATGTGTCACACAAGAAGCTGCTACAGAGGCTCATTACCAATGCCTATGACCA  
AACCTTGAGAGGGGATTATGTAGATATTACGACAGGTGAGCCACTTTTCCAAGGATAAGTTGCTTC  
GGTTGTTGGCCAAGTTAGCGTCCGATTCCAAGAGTGTGATTCAATTACAAGGATCTGAGCCATGGAA  
TGAGCGAATTGAAGTTCGTTCTGTCAGGCACTGCTACTGGTCATGTTCACAGATGGACCGCGGGAGTT  
AGCGGCCCTCCGTTACTGTATCAATTCTGCTCTTACGCTTGTGCCAGGATGAGATGAAAAAGCAGGATAT  
GGCTATCTATTGCTTACTTAAACAAATAA

20 MAEIYLAGGFWGLEYFSRISGVLETSGVYANGQVETNYQLLKETDHAETVQVIYDEKEVSLREILLYYFRVIDPLSI  
NQGNDRGRQYRTGIYYQDEADLPAYTVVQEQRMLGRKIAVEVEQLRHYIADYHQDYLRKNPSGYCHIDVTD  
DKPLIDAANYEKPSQEVLKASLSEESYRVTQEAATEAPFTNAYDQTFFEEGIYVDITGEPLFFAKDKFASCGWPSRPI  
SKELIHYYKDLSHGMERIEVRSRSGSAHLGHVFTDPRELGLLRYCINSASLRFVAKDEMEKAGYGYLLPYLNKZ

25 ID17 870bp

ATGAAGATTATTGTACCTGCAACCAAGTGCAATATCGGGCCAGGTTTACTCGGTGGGTAGCTGTAACCAAGT  
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ACGAGCGTAATCTTGTCTAAATCGCTTGCCTGAAATTGAGCTCAGACTGCAACCAAGACGCTTGAAGGAT  
TGATGTCCCTTGGCGCGCGTTGGTTCTTCAGCTCGTTATCGTGGATTGAACTAGCAACCAACTG  
GGTCAACTCAACTTATCAGACCATGAAAAATTGAGCTTGTGACCAAGATTGAAGGGCATCCTGACAATGTGGCT  
CCAGCATTATGGTAACTCGTTATGCAAGTTCTGTTGAAGGGCAAGTCTCTGCTATCGTAGCAGACTTCCAG  
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GTCTTATAAGGAAGCTGCTGCAAGTTCTATGCCAATGTAGCGTTGCTGCCCTGGTCAAGGAGACATGGT  
ACCGCTGGCAAGCAATCGAGGGAGACCTTCCATGAGCGCTATCGTCAAGGACTGGTAAGAGAATTGGCATG  
ATTAAGCAAGTGAACAAAGAAAATGGGCCATGCAACCTACCTTCTGGTCTGGCCAGCAGTTATGGTCTG  
GCTTCTCATGACAAGATGCCAACATAAGGAGAATTGAGAACCTTCAAGGAAACTGCAACCTTCAAAGGAAACTGCATGACTTG  
AGAGTTGATACCAAGGTGTCGTTAGAAGCAAAATAA

40 MKIIVPATSANIGPGFDSSVGAVTKYLQIEVCEERDEWLEHIQKWIPIHDERNLLKIALQIVPDLQPRRLKMTSDVPLA  
RGLGSSSSVIVAGIELANQLQLNLSDHEKLQLATKIEGHPDNVAPAIYGNLVIASSVEQVSAIVADFPECDFLAYIPNY  
ELRTRDSRSVPKKLSYKEAVAASSIANVAVAALLAGDMVTAGQAIEGDLFHERYRQDLDVREFAMIKQVTENGAYAT  
YLSGAGPTVMVLASHDKMPTIKAELEKQPFKGKLHDLRVDTQGVRVEAKZ

## 45

ID20 564bp

ATGAAATATCACGATTACATCTGGGATTAGGTGGAACTTACTGGATAATTATGAAACTTCAACAGCTGCATTG  
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TGCATTGAGACATTGCTCCAAATTAGAGAATTAGAAAAGTACAAGGAAATGAAGCCAGAGAGCTG  
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ACACCCGATTATTTGAAGGAATTCTGACTTATGGAAAGACATTCAAAATCAAGTGGCCGTCAATTGGTCT  
CTCATGCAATGATCAGGTTTGGAAATTAGAAAACCTCTATAGCAGCTTATTACAGAAGTGGTACTTC  
TAGCTCAGGTTTAAGGAAAGCCAATCCGAATCCATGCTTATTAAAGAGAAAAGTACAGATTAGCTCTGGT  
CTTGTCTTGTGATCGGCCGATTGATATCGAAGCAGGTCAAGCTGCAGGACTGATACCCACTTGTCTTACAGTA  
TCGTGAATTAAAGACAAGTATTAGACATATAA

## 55

MKYHDYIWBLGGTLLDNYETSTAATFETLALYGITQDHDSVYQALKVSTPFAIETFAPNLENFLCKENARELEHPI  
LFEGVSDLLEDISNQGGRHFLVSHRNDQVLEILEKTSIAAYTEVVTSSSGFKRKPNPESMLYREKYQISSGLVIGDRPID  
IEAGQAAGLDTHLFTSIVNLRQVL DIZ

## 60

ID21 1875bp  
ATGACAGAAGAAATCAAAATCTGCAGGCACAGGATTATGATGCCAGTCAAATTCAAGTTAGAGGGCTTAGAG  
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TTGATAACTCAATTGACGAGGCCTGGCAGGATTGCCAGCCATATTCAAGTTTATTGAGCCAGATGATTCGAT  
 TACTGTTGTGGATGATGGCGTGGTATCCCAGTCGATATTCAAGGAAAAACAGGCCGCTCTGCTGTTGAGACCGTC  
 TTACAGTCCTCACGCTGGAGAAAGTTCGGCGGTGGATACAAGGTTCTAGGTGGTCTTCACGGGGTGGGGT  
 CGTCAGTAGTTAATGCCCTTCACTCAATTAGACGTTCATGTTCAAAAATGGTAAGATTCACTTACAGAAGATA  
 CCGCGTGGTATGTTGTCGAGATCTGAAATAGTGGAGATACGGATAAAACAGGAACAACGTGTCACCTCACA  
 CCGGACCCAAAAATCTTCACTGAAACAACAATCTTCAAGGTTGATAAATTAACGGGATTCAGAGTGGCT  
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 AGGTGGGATTGCTAGTTACGTTGAATATACAGAAGGACAAACAGGATGTAATCTTGTACACCAATCTACAGAC  
 GGTGAGATGGATGATATACAGTTGAGGTAGCCATGCACTGACACAACGGTTACCATGAAAATGTCATGAGTTTC  
 10 GCCAATAATATTCAACCATGAAAGTGGAACACATGAACAAGGTTCCGTACAGCCTTGACACGTGTTATCAAC  
 GATTATGCTCGTAAAATAAGTTACTGAAAGACAATGAAGATAATTAAACAGGGGAAGATGTTGCGAAGGCTTA  
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 15 AATCTGGTTGGAAATTCCAAACCTTCCAGGGAAACTAGCAGACTGTTCTCTAATAACCCCTGCTGAAACAGAACT  
 CTTCATCGTCAAGGAGACTCAGCTGGTGGATCAGCCAAATCTGGTGTGAAACCGTGGTCTACGGCTATCCTCCA  
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 TCACAGCCATGGAACAGGATTGGCGCAGAATTGATGTTGAAAGCCGTTACCAAAAACCTGTTGATGAC  
 20 CGATGCCGATGTCGATGGAGCCCACATTCTGACCCCTTCTTAACTTGTGATTTATGAAACCAATCCTA  
 GAAGCTGGTTATGTTTATGCCCCAACCAACCTATGTTGTCAGGTTGGGAAGCGAGATAAGAATATTC  
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 TTCAGCTTATAAGGGCTAGGTGAAATGGACGATCATCAGTGTGGAAAACCATGGATCCGAACATCGCT  
 TGATGGCTAGAGTTCTGTAGATGATGTCAGAAGCAGATAAAATCTTGTATGTTGA  
 25 MTEEIKNLQAQDYDASQIQVLEGLEAVRMRPGMYIGSTSKEGLHHLVWEIVDNSIDEALAGFASHIQVFIEPDDSIIVVD  
 DGRGPVDIQEKTGRPAVETVFTVLHAGGKFGGGYKVSGGLHVGSSVNLSTQLDVHVHKNGKIHYQEYRRGHV  
 VADLEVGDTDKTGTGTVHFTPDPKIFTETTIFDFDKLNKRQLEFLNRLQISITDKRQGLEQTKHLYEGGLASYVEYI  
 NENKDVIIDPTYTDGEMDDITVEAMQYTTGYHENVMFSANNIHTHEQGFRRTALTRVINDYARKNKLKDNE  
 30 EDNLTGEDVREGLTAVISVKHPNPQFEGQTKLGNSEVVKTNRLFSEAFSDFLMENPQIAKRIVEKGILAAKARVAAK  
 RAREVTRKKSGLSINLPKGKADCSNNPAETELFIVEGDSAGGSAKSGRNREFQAILPRKGILNVEKASMDKILANEEIR  
 SLFTAMGTGFAEFDVSKARYQKLVLMTDADVDGAHIRTLLTLIYRMPKPILEAGYYVIAQPPIYGVKVGSEIKEYIQP  
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**ID54 1446bp**

35 ATAGTAGACGTTAAAAAAATCACGTTCACAGAAAGTGAAGCGAAGTGTAAATATAGTTGCTGACTATTATT  
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 40 AAATGCGACTCTAATTACTCAGAACATTCAATCAGTGTGCTGTTAGCAGATGTGAGATCGAAAATGTTACG  
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 45 TGTAGTGGATTGACACCTATGGCTCTATTAGTTCGGTGTGCGATCAGATGTCAACATCTGTGACTGTCAAT  
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 50 AGGCTCTCGGTTCTCTGAGCGCTACTCCTCTAGCAGATGGCAGTCAGGCCGGCGCAGTCAGAAAAGG  
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 55 TGA  
 MSRRFKKSRSQVKRSVNVLELTIYLVCFLFLFKYNILAFRFLNLYVLTALVLEVALVGLLIIYKKAEKFTIFLVEFSI  
 LVSSVSLFAVQQFVGLTNRLNATSNYSEYSISVAVLADSEJENVTQLTSVAPTSFENENIQKLLADIKSSQNPELTVNOS  
 SSYLAAYKSLIAGETKAIVLNSVFENIISEYSPDYASKIKIYTGKFTKKVEAPKTSKSQSFNIYVSGIDTYGPISSVSRSDVN  
 60 ILMTVNRDTKILLTTTPRDAYVPIADGGNNQKDKLTHAGIYGVDSIHLNLYGVLDINYYVRLNFFSELKIJDLGGI  
 DVYNDQEETAHNGKYYPAGNVHLDSEQAIIGFVRERYSLADGDRDRGRHQQKVIVAHQKLTSTEVEKNYSTIINSQ  
 DSQTNMPLMELMINLVNAOLESGGNYKVNSQDLKGTGRMDLPSYAMPDSNLYVMEIDDSSLAVYKAAIQDVMEGRZ

ID55 732bp

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 5 TGGCAGAACCTACAGACAGGGGGTGCAGAACCATTTCTACCGTGGAAATAGCTAAGGAAGTGGCGAGTGACTTGGTATTGC  
 GGAAGAGAAAGATAGCAGAAAACCTTCTACAGCTGGATAAGCTGGAAAAAGCGGATTCCGACCCCTCAATGA  
 TTACGGGGCTGAAATTATTACACACCAGATGTTCTGGATAAGCTGGAAAAAGCGGATTCCGACCCCTCAATGA  
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 10 AACTGATCGATATGGCTGTTACCGCAAGTAATAGTTCACATGTCCTCAACACCAAAACTTTGGCGAACGTTA  
 TAAATTATGAAAAAAAGAGCTAGTATTTAGAGCAGGATTGGTCATTGCAAGTGATATGACAAT  
 CTAGACGGTAGACCTCCATATGCCAGAACATGACCTGTTACCCAAAATACGGAGAACGAGGCTCAG  
 GAACCTTTATAGACAATCCTCGAAAAATTGATGCAACTAATTAG

15 MEDIHSIVFDVDDGPKSREESKALLAESYRQGVRTIVSTSHRKGMFETPEEKIAENFLQVREIAKEVASDLVIAYGAEI  
 YYTPDVLDKLEKKRIPTLNDSRYALIEFSMNTPYRDIHSALSKILMLGITPVIAHIERYDALENNEKRVRELIDMGCYTQV  
 NSSHVLKPILFGERYKFMKKRAQYFLEQDLVHVIASDMHNLDGRPHMAEAYDLVTQKYGEAKAQELFIDNPRKIVM  
 DQLIZ

ID58 3990bp

20 TTGATTTATATAATCGCTATCAATATAACAATGCAATCAGGAGGTTTGCAATGAAACATGAAAAACAAACAGCGTT  
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 25 ACTGATAAGGTAGCTAGTCTCCAAAACAGAAGAAAACACAAGAGGAAGTTAGTCAACTCCTAGTGTATAAA  
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 30 ATAAAGCGAAACATATTGCTACACTGATTTACACCTATTAGTCGGAAATGATGGACTCCGTTTATGTTGAGC  
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 AGGTATCGGCTCATTGACAGTAAATAGTCTGGACACATGGATGCGATTCTCAATGCCATGAAAGAATTGGG  
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 40 GTTCTATGTTGACTGGTGGAGGCTACGATGTCGCTCTCTAAACTACTAGCTGAAAAGGTACCCAAA  
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 45 TTGCAAAATGCCAACCGTGAATACTCGCAGCTGATTATGAAATCTGCAAGAGCAACACTTAAACGAGGTACCCAAA  
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 55 ATACATGGACTTCTGCAAGTAAAAGAATTCTCAACTTGGTACTGACGAATACGCCAACGATGCGACTAGT  
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 60 TGGTTCTCAAGAACAGCTATTGAGAAATACTGGAAAACACCATTCATCAACTAGCTTCTACCGTAAATATCCTGAA  
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 CTCTAAATTACAACCTCAACCGTAATAAACAGCTGAGGCTTGACACGCCCTGAGCAGCTAAACAG  
 AGGCCTCAACCGAGCTGTAACCTCATTCAGGAAGCCTAGATGAAAATGAAAGTGGCTGCCAATGTTGAAACAGACC  
 AGAACCTCATCACAAGAACTGAAGAAATTCCATTGAGATTATCAAGAAAGAAATCTAACCTCCAGGCCGTCA

GGAAAATATTACACAGCAGGAGTCAAAGGTGAACGAACCTCATTACATCTCTGTAECTCACTGAAAATGGAAAAAC  
 AACAGAAACAGTCCTTGATGCCAGGTAAACCAAGAAGTTATAAACCAAGTGGTGAAGTTGGCGCTCTGTAAC  
 TCACAAGGGTGTGAAAGTGGTCTTGCACCAACTACTGAGGTTAAACCTAGACTGGATATCCAAGAAGAAGAAAAT  
 TCCATTACACAGTGACTTGTGAAAATCCACTCTTACTCAAAGGAAAACACAAGTCACTACTAAGGGCGTCAAT  
 5 GGACATCGTAGCACTTCACTCTGTGAGCACCTCTGCCGATGGTAAGGAAGTGAACACTTGTAAATAGTGTGCG  
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 CGCCTATTGCTGAAGAAAAACCAAACAGAATCCCAAGCCACCAGCTCCATCAACTGCTCTGCTGAGGAAA  
 GCAAAGTTCTCCTCAAGATCCAGCCTGTGTTAAGAGAAAAACTTCCCTGAAACAGGAACTCACGATTCTG  
 10 CAGGACTAGTAGTCGAGGACTCATGCCAACACTAGCAGCTATGGACTCACTAAAAGAAAAGAAGACTAA  
 MIYIIAINITMQSGGFAMKHEKQQRFSIRKYAVGAASVLIGFAFQAQTVAADGVPTTENQPTIHTVSDSPQSSENRTEE  
 TPKAVLQPEAPKTVEETPATDKVASLPKTEEKPKQEEVSSTPSDKAEVVTPSAEKETANKKAEAPKKEEAKEVDSKE  
 SNTDKTDKDKPAKKDEAKAEADKPATEAGKERAATVNNEKLAKKKIVSIDAGRKYFSPQLKEIIDAKHYGYTDLHLL  
 15 VGNGLRFMLDDMSITANGTYASDDVKRAIEKGTDNYNDPNGNHLTESQMTDLINYAKDKGIGLIPTVNSPGHMD  
 AILNAMKELGIQNPNFSYFGKKSARTVLDNEQAVAFTKALIDKYAAYFAKTEIFNGLDEYANDATDAKGWSVLQA  
 DKYYPNEGYPVKGYEKFAYANDLARIVKSHGLKPMFDGYYNSDTSFGSFKDIVSMWTGGWGGYDVASSKLLA  
 EKGHQILNTNDAWYYVLGRNADGQGWYLDQGLNGIKNTPITSVPKTEGADIPPIGMVAAWADTPSARYSPSLFKL  
 MRHFANANAEEYPAADYESAEQALNEVPKDLNRYTAESVTAVKEAEKAIRSLDSNLRAQQDTIDQAIKLQETVNNLT  
 20 LTPEAQKEEEAKREVEKLAKNKVISIDAGRKYFTLNQLKRIVDKASELGYSVDVHLLGNDGLRFLDDMTITANGKTYA  
 SDDVKKAIIEGTKAYYDPNGTALTQAEVTELIEYAKSKDIGINPAINSPGHMDAMLVAMEKLGKNPQAHFDKVSKT  
 MDLKNEEAMNFVKALIGKYMDFLAGTKIFNFGTDEYANDATSAQGWYYLWKWYQLYGKFAEYANTLAAMAKERGL  
 QPMAFNDGFYEDKDDVQFDKDVLSYWSKGWYGYNLASPQYLASKGYKFLNTNGDWYILGQKPEDGGFLKKAI  
 ENTGKTPFNQLASTKYPEVDLPTVGSMSIADRSPEAYKEEIEFELMTAFADHNKDYFRANYNALREELAKIPTNLEG  
 25 YSKESLEALDAAKTALNYLNRNKQAEELDTLVANLKAALQGLKPAVTHSGSLDENEAANVETRPELITRTEIIPFEVI  
 KKENVLPNLPAGQENIIATGVKGERTHYISVLTENGTTETVLDSSQVTKEVINQVVEVGAPVTHKGDESGLAPTTEVKPR  
 DIQEEEIPFTTVTCENPLLKGKTQVITKGVNGHRNSFVSTSADGKEVKTIVNSVVAQEAVTQIVEVGMVTHVGDE  
 NGQAAIAEKKPLIPSQAPSTAPAEKSVLPQDPAPVVTEKKLPETGTHDSLGLVAGLMSTLAAYGLTKRKEDZ

ID122 825bp

30 ATGAACAAAAAAACAAGACAGACACTAATCGGACTGCTAGTGTATTGCTTGTACAGGGAGCTATTATATCA  
 AGCAGATGCCGTCGGCACCTAATAGTCCAAAACCAATCTAGTCAGAAAAAAACAAGCGTCTGAAGCTCTAGTC  
 AAGCATTGGCAGAGAGTGTCTTAACAGACGCAGTCAGAGTCAGCAATAAAGGGAGTCTGGAGTGAATGGCTCAG  
 GTGCTTTATCGTCATGGTAATAAAACAAATCTAGATGCCAAGCTAGTCAGTAAGCCCTACCGCTGACAATAAAAC  
 35 AAAGACAGTGGCAAGGAAACTGTCTCAACCTCTGGACTCCTCCAGGTTGGCATCAGGTCAAGAATCTAAAGGGCTCT  
 TCGTAAGAAACTGGGAATGGTTCAACTCTGGACTCCTCCAGGTTGGCATCAGGTCAAGAATCTAAAGGGCTCT  
 TATACCCATCGACTCGATAGAGGTCAATTGTTAGGCTATGCCCTAATCGGTGGTTGGATGGTTGATGCCCTCAA  
 CAAGCAATCTAAAACATTGCTGTCAGACAGCCTGGCAAATCAGGCACAAGCGAGTATTCGACTGGCTAAA  
 40 ACTACTATGAAAGCAAGGTGCGTAAAGCCTTGGACCAAAACAAGCGTGTCCGTTACCGTGTAAACCCTTACTACG  
 CTCAAACGAGGATTTAGTCCCTCAGCTCACAGATTGAAGCCAAGTCTCGGATGGAGAATTGGAAATTCAATGT  
 TCTAGTCCCAATGTTAAAAGGGACTCAACTGGATTACGAACTGGAGAAGTAACTGTAACTCACTA  
 MNKKTRQTLIGLLVLLLSTGSYYIKQMPSÄPNSPKTNLSQQKQASEÄPSQALEÄSVLTDÄVKSQIKGSLEWÑGSGAFIV  
 45 NGKTNLDAKVSSKPYADNKTKTGKETVPTVANALLSKATRQYKRNKETGNGSTWTPPGWHQVKNLKGSYTHAV  
 DRGHILLGYALIGGLDGFDASTSNPKNIAVQTAWANQQAQEYSTGQNYYESKVRKAIDQNKRVRYRVTLYYASNEDLV  
 PSASQIEAKSSDGELEFNVLVPNVQKGLQDRTGEVTQZ

ID123 225bp

50 GTGCTAACGATTAGCGGATTGAGGCAACTGATGAAGATGAATAAGAAATCAAGCTACGTAGTCAGGGTTACTT  
 TTAGTCATCATAGTACTGATTAGTACTCTGGCTCTAGGAATGGTTATGGTAGGTATGGAATCTGGGCA  
 AGGGTCAAGATCCATGGCTATCCTGCTCCAGCAAAATGGCAGGAATTGATTCATAAAATTACAGGAATTAG  
 VLRFSGERQVMKMNKKSSYVVKRLLVHVILGTLAEGIGEMVGYGLGKGQDPWAISPAKWQELIHKFTGNZ  
 55

## CLAIMS:

1. A *Streptococcus pneumoniae* protein or polypeptide having a sequence selected from those shown in table 1.
- 5 2. A *Streptococcus pneumoniae* protein or polypeptide having a sequence selected from those shown in table 2.
3. A protein or polypeptide as claimed in claim 1 or claim 2 provided in  
10 substantially pure form.
4. A protein or polypeptide which is substantially identical to one defined in any one of claims 1 to 3.
- 15 5. A homologue or derivative of a protein or polypeptide as defined in any one of claims 1 to 4.
6. An antigenic and/or immunogenic fragment of a protein or polypeptide as defined in Tables 1-3.
- 20 7. A nucleic acid molecule comprising or consisting of a sequence which is:
  - (i) any of the DNA sequences set out in Table 1 or their RNA equivalents;
  - 25 (ii) a sequence which is complementary to any of the sequences of (i);
  - (iii) a sequence which codes for the same protein or polypeptide, as those sequences of (i) or (ii);

- (iv) a sequence which is substantially identical with any of those of (i), (ii) and (iii);
- 5 (v) a sequence which codes for a homologue, derivative or fragment of a protein as defined in Table 1.
8. A nucleic acid molecule comprising or consisting of a sequence which is:
- 10 (i) any of the DNA sequences set out in Table 2 or their RNA equivalents;
- (ii) a sequence which is complementary to any of the sequences of (i);
- (iii) a sequence which codes for the same protein or polypeptide, as those sequences of (i) or (ii);
- 15 (iv) a sequence which is substantially identical with any of those of (i), (ii) and (iii);
- (v) a sequence which codes for a homologue, derivative or fragment of a protein as defined in Table 2;
- 20 9. The use of a protein or polypeptide having a sequence selected from those shown in Tables 1-3, or homologues, derivatives and/or fragments thereof, as an immunogen and/or antigen.
- 25 10. An immunogenic and/or antigenic composition comprising one or more proteins or polypeptides selected from those whose sequences are shown in Tables 1-3, or homologues or derivatives thereof, and/or fragments of any of these.
- 30 11. An immunogenic and/or antigenic composition as claimed in claim 10 which is

a vaccine or is for use in a diagnostic assay.

12. A vaccine as claimed in claim 11 which comprises one or more additional components selected from excipients, diluents, adjuvants or the like.

5

13. A vaccine composition comprising one or more nucleic acid sequences as defined in Tables 1-3.

14. A method for the detection/diagnosis of *S.pneumoniae* which comprises the 10 step of bringing into contact a sample to be tested with at least one protein or polypeptide as defined in Tables 1-3, or homologue, derivative or fragment thereof.

15. An antibody capable of binding to a protein or polypeptide as defined in Tables 1-3, or for a homologue, derivative or fragment thereof.

15

16. An antibody as defined in claim 15 which is a monoclonal antibody.

17. A method for the detection/diagnosis of *S.pneumoniae* which comprises the step 20 of bringing into contact a sample to be tested and at least one antibody as defined in claim 15 or claim 16.

18. A method for the detection/diagnosis of *S.pneumoniae* which comprises the step of bringing into contact a sample to be tested with at least one nucleic acid sequence as defined in claim 7 or claim 8.

25

19. A method of determining whether a protein or polypeptide as defined in Tables 1-3 represents a potential anti-microbial target which comprises inactivating said protein or polypeptide and determining whether *S.pneumoniae* is still viable.

30

20. The use of an agent capable of antagonising, inhibiting or otherwise interfering

with the function or expression of a protein or polypeptide as defined in Tables 1-3 in the manufacture of a medicament for use in the treatment or prophylaxis of *S.pneumoniae* infection

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